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OM protein - protein search, using sw model

Run on: December 19, 2003, 15:29:12; Search time 48 Seconds

(without alignments)

3845.813 Million cell updates/sec

Title: US-09-830-972-2

Perfect score: 5848

1 MEDIDQSSLVSSSTDSPPRP.....VKDAMAKIQAKIPGLKRKAD 1163 Sequence:

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 1107863 segs, 158726573 residues

Total number of hits satisfying chosen parameters: 1107863

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : A Geneseq 19Jun03:*

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/SIDS1/gcgdata/geneseq/geneseqp-emb1/AA2002.DAT:* 23: /SIDS1/qcqdata/qeneseq/qeneseqp-emb1/AA2003.DAT:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed,

SUMMARIES

		%				
Result	_	Query				December
No.	Score	Match	Length	DB 	ID	Description
1	5848	100.0	1163	23	ABB81074	Rat neurotransmitt
2	5846	100.0	1163	21	AAY71310	Rat neurite growth
3	5840	99.9	1162	21	AAY71557	Rat Nogo A truncat
4	5823	99.6	1163	21	AAY71384	Alternative versio
5	4921	84.1	974	21	AAY71560	Rat Nogo A protein
6	4403.5	75.3	1192	22	AAU04591	Human Nogo protein
7	4403.5	75.3	1192	23	ABP68600	Human pancreatic c
8	4398.5	75.2	1192	21	AAY56967	Human MAGI polypep
9	4398.5	75.2	1192	22	AAB82349	Human NOGO-A prote
10	4398.5	75.2	1192	23	ABB81078	Human neurotransmi
11	4398.5	75.2	1192	23	ABG30938	Human NogoA protei
12	4276.5	73.1	1178	21	AAY71311	Human neurite grow
13	4116	70.4	1246	22	AAU33228	Novel human secret
14	4023	68.8	803	21	AAY71562	Rat Nogo A protein
15	3714	63.5	737	21	AAY71386	Rat Nogo A protein
16	3699.5	63.3	746	21	AAY71391	Rat Nogo A protein
17	3651.5	62.4	736	21	AAY71398	Rat Nogo A protein
18	3630.5	62.1	732	21		Rat Nogo A protein
19	3494	59.7	695	21		Rat Nogo A protein
20	3436	58.8	684	21		Rat Nogo A protein
21	3385.5	57.9	983	24		Human MDDT polypep
22	3280.5	56.1	893	21		Human secreted pro
23	2779	47.5	552	21		Rat Nogo A protein
24	2500.5	42.8	642	19		Human secreted pro
25	2500.5	42.8	642	22		Human BG160_1 prot
26	2432	41.6	502	21		Rat Nogo A protein
27	2388	40.8	475			Rat Nogo A protein
28	2291	39.2	457			Rat Nogo A protein
29	1987	34.0	403	21		Rat Nogo A protein
- 30	1868	31.9	417			Rat Nogo A protein
31	1801	30.8	356			Rat Nogo A protein
32	1795.5	30.7	374			Rat Nogo A protein
33	1416	24.2	361			Alternative versio
34	1411.5	24.1	360			Rat neurite growth
35	1411.5	24.1	360			Rat neurotransmitt
36	1405.5		359			Rat Nogo A protein
37	1191	20.4	373			Human Nogo B prote
38	1191	20.4	373			Human MAGI polypep
39	1191	20.4	373			A bone marrow secr
40	1191	20.4	373			Human NOGO-B prote
41	1191	20.4	373			Human pancreatic c
42	1191	20.4	373	23		Human neurotransmi
43	1191	20.4	373	23		Human RTN4B SEQ ID
44	1183	20.2	373			Human NogoB protei
45	1063	18.2	284	21	AAY95030	Human clone vb22_1

```
RESULT 1
ABB81074
     ABB81074 standard; Protein; 1163 AA.
XX
AC
     ABB81074;
XX
DT
     05-NOV-2002 (first entry)
XX
DE
     Rat neurotransmitter receptor protein Nogo-A.
XX
KW
     Nerve regeneration; neuroprotection; neuronal degeneration; CNS; PNS;
KW
     central nervous system; peripheral nervous system; tranquillizer; Nogo;
     vulnerary; cerebroprotective; anti-tumour; antidiabetic; anticonvulsant;
KW
KW
     nootropic; antiparkinsonian; ophthalmological; analgesic; hepatotropic;
KW
     osteopathic; vasotropic; nephrotropic; cytostatic; antigen; gene therapy;
KW
     neurotransmitter receptor; rat; receptor.
XX
OS
     Rattus norvegicus.
XX
PN
     US2002072493-A1.
XX
PD
     13-JUN-2002.
XX
PF
     28-JUN-2001; 2001US-0893348.
XX
PR
     19-MAY-1998;
                    98IL-0124500.
PR
     21-JUL-1998;
                    98WO-US14715.
PR
     22-DEC-1998;
                    98US-0218277.
PR
     19-MAY-1999;
                    99US-0314161.
XX
PΑ
     (YEDA ) YEDA RES & DEV CO LTD.
XX
PI
     Eisenbach-Schwartz M, Hauben E, Cohen IR, Beserman P, Mosonego A;
PΙ
     Moalem G;
XX
DR
     WPI; 2002-607255/65.
DR
     N-PSDB; ABN86600.
XX
PT
     Promoting nerve regeneration and preventing neuronal degeneration in
PT
     the central/peripheral nervous system from injury/disease, comprises
PT
     administering nervous system-specific activated T cells/antigen, or
PT
     analogs/peptides
XX
PS
     Example 5; Page 44-47; 93pp; English.
XX
CC
     The invention relates to promoting nerve regeneration or conferring
CC
     neuroprotection and preventing or inhibiting neuronal degeneration in the
CC
     central/peripheral nervous system (NS). The method involves administering
CC
      NS-specific activated T cells, NS-specific antigen, its analogue or its
CC
     peptide, a nucleotide sequence the NS-specific antigen or its analogue or
CC
     combinations. The method is useful for promoting nerve regeneration and
CC
     preventing neuronal degeneration in central/peripheral nervous system
CC
     from injury/disease, where the injury is spinal cord injury, blunt
CC
     trauma, penetrating trauma, hemorrhagic stroke, ischaemic stroke or
CC
     damages caused by surgery such as tumour excision. The disease is not an
CC
     autoimmune disease or neoplasm. The disease results in a degenerative
CC
     process occurring in either gray or white matter or both. The disease
```

```
CC
    is diabetic neuropathy, senile dementia, Alzheimer's disease, Parkinson's
    disease, facial nerve (Bell's) palsy, glaucoma, Huntington's chorea,
CC
CC
    amyotrophic lateral sclerosis, non-arteritic optic neuropathy, and
    vitamin deficiency, intervertebral disc herniation, prion diseases such
CC
CC
    as Creutzfeldt-Jakob disease, carpal tunnel syndrome, peripheral
CC
    neuropathies associated with various diseases, including but not limited
CC
    to urėmia, porphyria, hypoglycemia, Sjorgren Larsson syndrome, acute
    sensory neuropathy, chronic ataxic neuropathy, biliary cirrhosis, primary
CC
    amyloidosis, obstructive lung diseases, acromegaly, malabsorption
CC
    syndromes, polycythemia vera, immunoglobulin (Ig)A- and IqG gamma-
CC
CC
    pathies, complications of various drugs (e.g., metronidazole) and toxins
CC
    (e.g., alcohol or organophosphates), Charcot-Marie-Tooth disease, ataxia
    telangectasia, Friedreich's ataxia, amyloid polyneuropathies,
CC
    adrenomyeloneuropathy, Giant axonal neuropathy, Refsum's disease, Fabry's
CC
    disease, or lipoproteinemia. The present sequence represents the rat
CC
    neurotransmitter receptor protein Nogo-A, an example of NS-specific
CC
CC
    antigen.
XX
SO
    Sequence
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 Query Match
                      100.0%; Score 5848; DB 23; Length 1163;
 Best Local Similarity
                      100.0%; Pred. No. 1.1e-296;
 Matches 1163; Conservative
                         0; Mismatches
                                           0; Indels
                                                       0; Gaps
                                                                 0;
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Qу
            Db
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Qу
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Dh
        121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
Qу
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Db
Qу
        181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240
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        181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240
        241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300
Qу
            Db
        241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300
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Qу
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Qу
        361 PVREEYADFKPFEOAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEOKSLGKDSEGR 420
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        421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480
Qу
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481 QIITEKTSPKTSNPFLVAVODSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVOEACESEL 540

Qу

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Db	541		600
Qу	601	LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF	660
Db	601		660
Qу	661	NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720
Db	661	NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720
Qу	721	PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQP	780
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Qу	781	NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	840
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Db	841	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK	900
Qу	901	DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED	960
Db	901	DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED	960
Qу	961	RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS	1020
Db	961	RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS	1020
Qу		VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	
Db	1021	VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	1080
Qу		FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLA	
Db		FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLA	1140
QУ		NKSVKĎAMAKIQAKI PGLKŘKÁĎ 1163	
Db	1141	NKSVKDAMAKIQAKIPGLKRKAD 1163	
	AAY71310 02-NOV-2		

```
XX
KW
     Rat; neurite growth inhibitor; Nogo A; neural cell; myelin; CNS;
     central nervous system; neoplastic disease; antiproliferative; glioma;
KW
     antisense qene therapy; neuroblastoma; menagioma; retinoblastoma;
     degenerative nerve disease; Alzheimer's disease; Parkinson's disease;
KW
     hyperproliferative disorder; benign dysproliferative disorder; diagnosis;
KW
KW
     psoriasis; tissue hypertrophy; neuronal regeneration; treatment;
     structural plasticity; screening.
KW
XX
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XX
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FT
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XX
PR
                    98US-0107446.
     06-NOV-1998;
XX
PΑ
     (SCHW/) SCHWAB M E.
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PΑ
    (CHEN/) CHEN M S.
XX
PΙ
    Schwab ME, Chen MS;
XX
    WPI; 2000-400052/34.
DR
    N-PSDB; AAD01173.
DR
XX
    Nogo proteins and nucleic acids useful for treating neoplastic
РΤ
    disorders of the central nervous system and inducing regeneration of
РΤ
PT
    neurons -
XX
    Claim 3; Fig 2A; 122pp; English.
PS
XX
    The present sequence is a rat Nogo A protein which is a
CC
    potent neural cell growth inhibitor and is free of all central nervous
CC
    system (CNS) myelin material with which it is natively associated. The
CC
    protein was derived from a cDNA generated by fusing RO18U37-3, R1-3U21
CC
    cDNAs isolated from hexanucleotides-primed rat brain stem/spinal cord
CC
    library, and Oli18 cDNA from an oligo d(T)-primed rat oligodendrocyte
CC
    library. Nogo proteins and fragments displaying neurite growth
CC
    inhibitory activity are used in the treatment of neoplastic disease of
CC
    the CNS e.g. glioma, glioblastoma, medulloblastoma, craniopharyngioma,
CC
    ependyoma, pinealoma, haemangioblastoma, acoustic neuroma,
CC
    oligodendroglioma, menagioma, neuroblastoma or retinoblastoma and
CC
    degenerative nerve diseases e.g. Alzheimer's and Parkinson's diseases.
CC
    Therapeutics which promote Nogo activity can be used to treat or prevent
CC
    hyperproliferative or benign dysproliferative disorders e.g. psoriasis
CC
    and tissue hypertrophy. Ribozymes or antisense Nogo nucleic acids can be
CC
    used to inhibit production of Nogo protein to induce regeneration of
CC
    neurons or to promote structural plasticity of the CNS in disorders where
CC
    neurite growth, regeneration or maintenance are deficient or desired.
CC
    The animal models can be used in diagnostic and screening methods for
CC
    predisposition to disorders and to screen for or test molecules which
CC
    can treat or prevent disorders or diseases of the CNS.
CC
    Note: The present sequence designated as SEQ ID NO: 2 is stated to
CC
    be the same as the sequence shown in Fig. 13 (see AAY71384) of the
CC
     specification. However, this sequence does not match the sequence given
CC
     in Fig. 13. SEQ ID numbers 35-42 are referred in claim 32 and
CC
CC
     SEQ ID NO: 29 in disclosure of the specification. However, the
     specification does not include sequences for these SEQ ID numbers.
CC
XX
SO
              1163 AA;
    Sequence
                         100.0%; Score 5846; DB 21; Length 1163;
  Query Match
                        99.9%; Pred. No. 1.4e-296;
  Best Local Similarity
                                                                 Gaps
                                                                         0;
  Matches 1162; Conservative 1; Mismatches
                                                0; Indels
           1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEDEEEDDEDLEELEVLERK 60
Qу
             1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
Db
          61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120
Qу
             61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120
Db
         121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
Qу
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Db	121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180	
Qу	181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240	
Db	181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240	
Qу	241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300	
Db	241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300	
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Db	301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVA 360	
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Db	361 PVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCIEDSLEQKSLGKDSEGR 420	
Qу	421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480	
Db	421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480	
Qy Db	481 QIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESEL 540	
Qу	541 NEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSL 600	
Db		
QУ	601 LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF 660	
Db		
Qу	661 NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE 720	
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Qу	721 PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQP 780	
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QУ	781 NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE 840	
Db	781 NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE 840	
Qу	841 IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK 900	
Db	841 IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK 900	
Qу	901 DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED 960	
Db	901 DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED 960	
Qy Db	961 RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS 1020 	
עע	NOT VORDWARDED VIOLAN AND MANAGEMENT AND MANAGEMENT TO TAKE THE PROPERTY AND TOTAL WORD AND THE PROPERTY TO A TAKE THE PROPERTY AND THE PROPER	

```
1021 VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL 1080
Qу
             Db
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RESULT 3
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XX
AC
    AAY71557;
XX
DT
    02-NOV-2000 (first entry)
XX
    Rat Nogo A truncated protein used in the construction of mutant Nogo-A.
DE
XX
    Rat; neurite growth inhibitor; Nogo A; neural cell; myelin; CNS;
KW
KW
    central nervous system; neoplastic disease; antiproliferative; glioma;
    antisense gene therapy; neuroblastoma; menagioma; retinoblastoma;
KW
    degenerative nerve disease; Alzheimer's disease; Parkinson's disease;
KW
    hyperproliferative disorder; benign dysproliferative disorder; diagnosis;
KW
    psoriasis; tissue hypertrophy; neuronal regeneration; treatment;
KW
    structural plasticity; screening; mutant; mutein.
KW
XX
OS
    Rattus sp.
XX
PN
    WO200031235-A2.
XX
PD
    02-JUN-2000.
XX
PF
    05-NOV-1999;
                  99WO-US26160.
XX
PR
    06-NOV-1998;
                  98US-0107446.
XX
PΑ
     (SCHW/) SCHWAB M E.
PΑ
     (CHEN/) CHEN M S.
XX
PΙ
    Schwab ME,
               Chen MS;
XX
DR
    WPI; 2000-400052/34.
XX
    Nogo proteins and nucleic acids useful for treating neoplastic
PТ
PT
    disorders of the central nervous system and inducing regeneration of
PT
    neurons -
XX
PS
    Example; Page -; 122pp; English.
XX
CC
    The patent relates to neurite growth inhibitor Nogo which is free of
CC
    all central nervous system (CNS) myelin material with which it is
```

CCnatively associated. Nogo proteins and fragments displaying neurite CC growth inhibitory activity are used in the treatment of neoplastic CC disease of the CNS e.g. glioma, glioblastoma, medulloblastoma, CC craniopharyngioma, ependyoma, pinealoma, haemangioblastoma, acoustic CCneuroma, oligodendroglioma, menagioma, neuroblastoma or retinoblastoma CC and degenerative nerve diseases e.g. Alzheimer's and Parkinson's CC diseases. Therapeutics which promote Nogo activity can be used to treat CC or prevent hyperproliferative or benign dysproliferative disorders e.g. CC psoriasis and tissue hypertrophy. Ribozymes or antisense Nogo nucleic CC acids can be used to inhibit production of Nogo protein to induce CC regeneration of neurons or to promote structural plasticity of the CNS CC in disorders where neurite growth, regeneration or maintenance are CCdeficient or desired. The animal models can be used in diagnostic and CC screening methods for predisposition to disorders and to screen for or CC test molecules which can treat or prevent disorders or diseases of the CCCNS. The present sequence is a truncated form of rat Noqo A protein shown in AAY71310, which is used in the construction of mutant Noqo-A. Noqo-A CC CC is composed of His-tag/T7-tag/vector/Nogo-A sequence aa 1-1162. CC Nogo A deletion mutants were used for mapping the inhibitory sites of CC Nogo protein. Major inhibitory region was identified in the CC Nogo A sequence from amino acids 172-974, particularly amino acids 542-722. In addition, N-terminal region 1-171 was found to be inhibitory CC CC to NIH 3T3 fibroblast spreading. CC Note: The present sequence is not given in the specification but is CC derived from rat Nogo A sequence shown in AAY71310. SEQ ID numbers 35-42 CC are referred in claim 32 and SEQ ID NO: 29 in disclosure of the CC specification. However, the specification does not include sequences for CC these SEQ ID numbers. XX SQ Sequence 1162 AA; Query Match 99.9%; Score 5840; DB 21; Length 1162; Best Local Similarity 99.9%; Pred. No. 2.8e-296; Matches 1161; Conservative 1; Mismatches 0; Indels 0; Gaps 0: 1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60 Qу Dh 1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60 61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120 Qу Db 61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120 121 PÄAAVLPSKLPEDDEPPARPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180 Qу Db 121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180 181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240 Qу 181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240 Db 241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300 Qу Db 241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300 301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVA 360 Qу

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Db	361	PVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCIEDSLEQKSLGKDSEGR	420
Qу	421	NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA	480
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Qу	601	LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF	660
Db	601	LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF	660
Qу	661	NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720
Db		NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720
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Db		NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	
Qу	•	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK	
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Db		RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	
Qy Db		VIISFRIINGVIQAIQKSDEGHPFRAILESEVAISEELVQKISNSALGHVNSIIKELRRL	
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Qy Db		FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLTLALISLFSTPVTTERHQVQTDHYLGLA	
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RESULT 4
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ID
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XX
AC
     AAY71384;
XX
DT
     02-NOV-2000
                 (first entry)
XX
DΕ
     Alternative version of rat neurite growth inhibitor Nogo A.
XX
KW
     Rat; neurite growth inhibitor; Nogo A; neural cell; myelin; CNS;
KW
     central nervous system; neoplastic disease; antiproliferative; qlioma;
KW
     antisense gene therapy; neuroblastoma; menagioma; retinoblastoma;
KW
     degenerative nerve disease; Alzheimer's disease; Parkinson's disease;
     hyperproliferative disorder; beniqn dysproliferative disorder; diagnosis;
KW
     psoriasis; tissue hypertrophy; neuronal regeneration; treatment;
KW
     structural plasticity; screening.
KW
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OS
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XX
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FT
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FT
FT
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FT/note= "There is Ile at this position in the FΤ sequence shown in AAY71310" FTMisc-difference 469 FТ /label= Unknown /note= "There is Lys at this position in the тч FTsequence shown in AAY71310" FTMisc-difference 661 /note= "There is Asn at this position in the FΤ FTsequence shown in AAY71310" FTMisc-difference 820 FT/note= "There is Leu at this position in the FTsequence shown in AAY71310" XXPNWO200031235-A2. XX PD 02-JUN-2000. XX 05-NOV-1999; PF 99WO-US26160. ХX PR 06-NOV-1998; 98US-0107446. XX PA (SCHW/) SCHWAB M E. (CHEN/) CHEN M S. $D\Delta$ XX PΙ Schwab ME, Chen MS; XX WPI; 2000-400052/34. DR XX PTNogo proteins and nucleic acids useful for treating neoplastic РТ disorders of the central nervous system and inducing regeneration of PTneurons -XX PS Claim 3; Fig 13; 122pp; English. XXCC The present sequence is an alternative version of rat Nogo A protein CC which is a potent neural cell growth inhibitor and is free of all CC central nervous system (CNS) myelin material with which it is CC natively associated. Nogo proteins and fragments displaying CC neurite growth inhibitory activity are used in the treatment of neoplastic disease of the CNS CC CC e.g. glioma, glioblastoma, medulloblastoma, craniopharyngioma, CC ependyoma, pinealoma, haemangioblastoma, acoustic neuroma, CC oligodendroglioma, menagioma, neuroblastoma or retinoblastoma and CC degenerative nerve diseases e.g. Alzheimer's and Parkinson's diseases. CC Therapeutics which promote Nogo activity can be used to treat or prevent CC hyperproliferative or benign dysproliferative disorders e.g. psoriasis CC and tissue hypertrophy. Ribozymes or antisense Nogo nucleic acids can be CC used to inhibit production of Nogo protein to induce regeneration of CC neurons or to promote structural plasticity of the CNS in disorders where CC neurite growth, regeneration or maintenance are deficient or desired. CC The animal models can be used in diagnostic and screening methods for CC predisposition to disorders and to screen for or test molecules which CC can treat or prevent disorders or diseases of the CNS. CC Note: The present sequence is an alternative version of the CC Nogo A sequence shown in Fig. 2A (see AAY71310). CCSEQ ID numbers 35-42 are referred in claim 32 and SEQ ID NO: 29 CC in disclosure of the specification. However the specification does not

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CC
    include sequences for these SEO ID numbers.
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SO
   Sequence
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 Query Match
                   99.6%;
                         Score 5823; DB 21;
                                         Length 1163;
 Best Local Similarity
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 Matches 1159; Conservative
                         0; Mismatches
                                         Indels
                                                   Gaps
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Qу
          Db
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
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Qу
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Db
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Qу
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Db
Qу
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           Db
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Qу
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Db
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Qу
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XX
AC
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XX
DT
    02-NOV-2000 (first entry)
XX
DE
    Rat Nogo A protein fragment used in the construction of mutant NiAext.
XX
KW
    Rat; neurite growth inhibitor; Nogo A; neural cell; myelin; CNS;
KW
    central nervous system; neoplastic disease; antiproliferative; glioma;
KW
    antisense gene therapy; neuroblastoma; menagioma; retinoblastoma;
KW
    degenerative nerve disease; Alzheimer's disease; Parkinson's disease;
KW
    hyperproliferative disorder; benign dysproliferative disorder; diagnosis;
KW
    psoriasis; tissue hypertrophy; neuronal regeneration; treatment;
KW
    structural plasticity; screening; mutant; mutein.
XX
O$
    Rattus sp.
XX
ΡN
    WO200031235-A2.
XX
PD
    02-JUN-2000.
XX
PF
    05-NOV-1999;
                99WO-US26160.
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XX
PR
     06-NOV-1998;
                    98US-0107446.
ХX
     (SCHW/) SCHWAB M E.
PΑ
     (CHEN/) CHEN M S.
PΑ
XX
PΙ
     Schwab ME,
                 Chen MS;
XX
DR
     WPI; 2000-400052/34.
XX
PT
     Nogo proteins and nucleic acids useful for treating neoplastic
PT
     disorders of the central nervous system and inducing regeneration of
PT
     neurons -
XX
PS
     Example; Page -; 122pp; English.
XX
CC
     The patent relates to neurite growth inhibitor Nogo which is free of
CC
     all central nervous system (CNS) myelin material with which it is
CC
     natively associated. Nogo proteins and fragments displaying neurite
CC
     growth inhibitory activity are used in the treatment of neoplastic
CC
     disease of the CNS e.g. glioma, glioblastoma, medulloblastoma,
CC
     craniopharyngioma, ependyoma, pinealoma, haemangioblastoma, acoustic
CC
     neuroma, oligodendroglioma, menagioma, neuroblastoma or retinoblastoma
CC
     and degenerative nerve diseases e.g. Alzheimer's and Parkinson's
CC
     diseases. Therapeutics which promote Nogo activity can be used to treat
CC
     or prevent hyperproliferative or benign dysproliferative disorders e.g.
CC
     psoriasis and tissue hypertrophy. Ribozymes or antisense Nogo nucleic
CC
     acids can be used to inhibit production of Nogo protein to induce
CC
     regeneration of neurons or to promote structural plasticity of the CNS
CC
     in disorders where neurite growth, regeneration or maintenance are
CC
     deficient or desired. The animal models can be used in diagnostic and
CC
     screening methods for predisposition to disorders and to screen for or
CC
     test molecules which can treat or prevent disorders or diseases of the
CC
     CNS. The present sequence is a fragment of rat Nogo A protein shown in
CC
     AAY71310, which is used in the construction of mutant NiAext. The mutant
CC
     is composed of His-tag/T7-tag/vector/Nogo-A sequence aa 1-974/T7-tag.
CC
     Nogo A deletion mutants were used for mapping the inhibitory sites of
CC
     Nogo protein. Major inhibitory region was identified in the
CC
     Nogo A sequence from amino acids 172-974, particularly amino acids
     542-722. In addition, N-terminal region 1-171 was found to be inhibitory
CC
     to NIH 3T3 fibroblast spreading.
CC
CC
     Note: The present sequence is not given in the specification but is
CC
     derived from rat Nogo A sequence shown in AAY71310. SEQ ID numbers 35-42
CC
     are referred in claim 32 and SEQ ID NO: 29 in disclosure of the
CC
     specification. However, the specification does not include sequences for
CC
     these SEQ ID numbers.
XX
SO
     Sequence
                974 AA;
  Query Match
                          84.1%;
                                 Score 4921; DB 21; Length 974;
  Best Local Similarity
                                 Pred. No. 1.9e-248;
                         99.9%;
  Matches 973; Conservative
                                1; Mismatches
                                                                            0;
                                                  0; Indels
                                                                    Gaps
Qy
            1 MEDIDOSSLVSSSTDSPPRPPPAFKYOFVTEPEDEEDEEEEEDEEDDEDLEELEVLERK 60
              Db
            1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
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Qy	61	PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP	120	
Db	61	PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP	120	
Qy	121	PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP	180	
Db	121	PAAAVLPSKLPEDDEPPARPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP	180	
Qy	181	AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL	240	
Db	181	AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL	240	
Qy	241	GNLSAVSSSEGTIEETLNEÄSKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES	300	
Db	241	GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES	300	
Qy	301	AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVA	360	
Db	301	AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVA	360	
Qу	361	PVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGR	420	
Db	361	PVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCIEDSLEQKSLGKDSEGR	420	
Qy	421	NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA	480	
Db	421	NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA	480	
Qy	481	QIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESEL	540	
Db	481	QIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESEL	540	
Qy	541	NEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSL	600	
Db	541	NEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSL	600	
Qy	601	LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF	660	
Db		LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF		
Qy	661	NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720	
Db	661	NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720	
Qу		PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQP		
Db		PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQP		
QУ	781	NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	840	
Db		NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE		
Qу	841	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK	900	
Db		IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK		
Qу	901	DEVHVSDEFSENRSSVSKAS I SPSNVSALEPOTEMGS I VKSKSLTKEAEKKLPSDTEKED	960	

```
Db
          901 DEVHVSDEFSENRSSVSKASISPSNVSALEPOTEMGSIVKSKSLTKEAEKKLPSDTEKED 960
          961 RSLSAVLSAELSKT 974
QУ
              111111111111
          961 RSLSAVLSAELSKT 974
Db
RESULT 6
AAU04591
     AAU04591 standard; Protein; 1192 AA.
XX
AC
    AAU04591;
XX
DT
     26-SEP-2001 (first entry)
XX
DE
     Human Nogo protein.
XX
KW
     Human; Nogo receptor; axonal growth; immunogen; antibody; nogo protein;
     cranial trauma; cerebral trauma; spinal cord injury; stroke;
KW
KW
     demyelinating disease; multiple sclerosis; monophasis demyelination;
KW
     encephalomyelitis; multifocal leukoencephalopathy; panencephalitis;
KW
     Marchiafava-Bignami disease; pontine myelinolysis; adrenoleukodystrophy;
KW
     Pelizaeus-Merzbacher disease; Spongy degeneration; Alexander's disease;
KW
     Canavan's disease; metachromatic leukodystrophy; viral infection;
ΚW
     Krabbe's disease.
XX
OS
     Homo sapiens.
XX
FΗ
     Key
                     Location/Qualifiers
FT
                     1054..1119
     Domain
FT
                     /label= Lumenal extracellular domain
                     /note= "This sequence is specifically claimed"
FT
FT
     Peptide
                     1055..1094
FT
                     /label= Pep1
FT
                     /note= "Receptor binding inhibitory peptide. This
FT
                     sequence is specifically claimed"
FT
     Peptide
                     1064..1088
FT
                     /label= Pep2
FT
                     /note= "Receptor binding inhibitory peptide. This
FT
                     sequence is specifically claimed"
FT
                     1074..1098
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                     /label= Pep3
FT
FΤ
                     /note= "Receptor binding inhibitory peptide. This
FT
                     sequence is specifically claimed"
FT
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                     1084..1108
FT
                     /label= Pep4
FT
                     /note= "Receptor binding inhibitory peptide. This
FΤ
                     sequence is specifically claimed"
                     1095..1119
FΤ
     Peptide
FT
                     /label= Pep5
FT
                     /note= "Receptor binding inhibitory peptide. This
FT
                    sequence is specifically claimed"
XX
PN
    WO200151520-A2.
XX
PD
    19-JUL-2001.
```

```
XX
PF
    12-JAN-2001; 2001WO-US01041.
XX
PR
    12-JAN-2000; 2000US-0175707.
PR
    26-MAY-2000; 2000US-0207366.
PR
    29-SEP-2000; 2000US-0236378.
ХX
PΑ
    (UYYA ) UNIV YALE.
XX
PΙ
    Strittmatter SM;
XX
DR
    WPI; 2001-442138/47.
    N-PSDB; AAS09453.
DR
XX
PT
    Novel Nogo receptor protein useful for identifying modulator of Nogo
PT
    protein or Nogo receptor protein, which is useful for treating central
PT
    nervous system disorders
XX
PS
    Example 1; Page 101-104; 109pp; English.
XX
CC
    The sequence is the human Nogo protein, a 250kDa myelin-associated axon
CC
    growth inhibitor. The invention relates to the use of the nogo receptor,
    nogo protein, their nucleic acids, vectors expressing them and antibodies
CC
CC
    against them, to isolate agents which block nogo receptor mediated axonal
CC
    growth. The agent is useful for treating a central nervous system
CC
    disorder which is a result of cranial or cerebral trauma, spinal cord
CC
    injury, stroke or a demyelinating disease selected from multiple
CC
    sclerosis, monophasis demyelination, encephalomyelitis, multifocal
CC
    leukoencephalopathy, panencephalitis, Marchiafava-Bignami disease,
CC
    pontine myelinolysis, adrenoleukodystrophy, Pelizaeus-Merzbacher disease,
CC
    Spongy degeneration, Alexander's disease, Canavan's disease,
CC
    metachromatic leukodystrophy, viral infection and Krabbe's disease.
XX
SO
    Sequence
              1192 AA;
 Query Match
                       75.3%; Score 4403.5; DB 22; Length 1192;
 Best Local Similarity
                       75.9%; Pred. No. 2.5e-221;
 Matches 909; Conservative 104; Mismatches 145; Indels
                                                         39; Gaps
                                                                    20:
          1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
QУ
            Db
          1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDEDLEELEVLERK 58
Qу
          61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
            Dh
         59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP 118
         116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
Qу
                  Db
         119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
         167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Oy
            Db
         179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEOPGNTISAGOEDFPSVLLETAASLPS 237
Qу
         226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
```

Db	238	LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY	297
Qy	286	SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVGKED	339
Db	298	SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNQQELPTALTKLVKED	357
Qу	340	RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANV	395
Db	358	EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL	416
Qу	396	ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT	454
Db [®]	417	ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRSGAYITCAPFNPAATESIA	476
Qy	455	ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS	513
Db	477	TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT	536
Qy	514	KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL	573
Db	537	KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL	596
Qу	574	CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN	633
Db	597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN	655
Qy	634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF	692
Db	656	PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF	715
Qy	693	SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS	752
Db	716	SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS	775
Qу	753	-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF :: ::: :	809
Db	776	FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLQMEEL	835
Qy	810	NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS : :	868
Db	836	STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS	895
Qy	869	DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	926
Db	896	HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	955
Qy	927	SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	986
Db	956	SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK	1015
Qy	987	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1046
Db	1016	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1075
Qy	1047	YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1106
Db	1076	YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1135

```
1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
Qу
              Db
        1136 NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1192
RESULT 7
ABP68600
    ABP68600 standard; Protein; 1192 AA.
XX
AC
    ABP68600;
XX
DT
    14-JAN-2003 (first entry)
XX
DE
    Human pancreatic cancer expressed protein SEQ ID NO 71.
XX
KW
    Human; pancreas; cancer; qene therapy; vaccine; immunostimulant;
KW
    cytostatic; tumour.
XX
OS
    Homo sapiens.
XX
PN
    WO200260317-A2.
XX
PD
    08-AUG-2002.
XX
PF
    30-JAN-2002; 2002WO-US02781.
XX
PR
    30-JAN-2001; 2001US-265305P.
PR
     31-JAN-2001; 2001US-265682P.
PR
     09-FEB-2001; 2001US-267568P.
PR
    21-MAR-2001; 2001US-278651P.
     28-APR-2001; 2001US-287112P.
PR
     16-MAY-2001; 2001US-291631P.
     12-JUL-2001; 2001US-305484P.
PR
     20-AUG-2001; 2001US-313999P.
PR
     27-NOV-2001; 2001US-333626P.
PR
XX
PA
     (CORI-) CORIXA CORP.
XX
    Benson DR, Kalos MD, Lodes MJ, Persing DH, Hepler WT, Jiang Y;
PΙ
XX
DR
    WPI; 2002-627435/67.
DR
    N-PSDB; ABV94680.
XX
PT
    New isolated polynucleotide and pancreatic tumor polypeptides, useful
PT
    for diagnosing, preventing and/or treating cancer, particularly
PT
    pancreatic cancer
XX
PS
    Claim 2; SEQ ID NO 71; 300pp + Sequence Listing; English.
XX
CC
    The invention relates to an isolated polynucleotide (I) comprising: (a)
CC
    any of a group of over 4000 nucleotide sequences (ABV94628-ABV99145);
CC
     (b) complements of (a); (c) sequences consisting of at least 20
CC
    contiguous residues of (a); (d) sequences that hybridize to (a), under
CC
    moderately stringent conditions; (e) sequences having at least 75% or 90%
CC
     identity to (a); or (f) degenerate variants of (a). Polypeptides
CC
     (ABP68596-ABP68637) encoded by (I) and oligonucleotide can be used to
```

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polynucleotides, antibodies, fusion proteins, T cell populations and
CC
CC
    antigen presenting cells expressing the polypeptide are useful in
    treating pancreatic cancer and stimulating an immune response. The
CC
    polynucleotides can be used as probes or primers for nucleic acid
CC
    hybridisation, in the design and preparation of ribozyme molecules for
CC
    inhibiting expression of the tumour polypeptides and proteins in the
CC
    tumour cells, in vaccines and for gene therapy.
CC
    Note: The sequence data for this patent did not form part of the printed
CC
    specification, but was obtained in electronic format directly from WIPO
CC
CC
    at ftp.wipo.int/pub/published pct sequences.
XX
SQ
    Sequence
            1192 AA;
 Query Match.
                     75.3%; Score 4403.5; DB 23; Length 1192;
                     75.9%; Pred. No. 2.5e-221;
 Best Local Similarity
 Matches 909; Conservative 104; Mismatches 145; Indels
                                                   39; Gaps
                                                             20;
         1 MEDIDOSSLVSSSTDSPPRPPPAFKYOFVTEPEDEEDEEEEEDEEDDEDLEELEVLERK 60
QУ
           1 MEDLDOSPLVSSS-DSPPRPOPAFKYOFVREPEDEE-EEEEEEEEDEDEDLEELEVLERK 58
Db
         61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPEROPSWERSPAA---P 115
Qу
           59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP 118
Db
Qу
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                119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
Dh
        167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Qу
           179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEOPGNTISAGOEDFPSVLLETAASLPS 237
Db
Qу
        226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
           238 LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY 297
Db
        286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPOESPVG----KED 339
Qу
           Db
        298 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNQQELPTALTKLVKED 357
        340 RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAA----RANV 395
Qу
            358 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 416
Db
        396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
Qу
           Db
        417 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRSGAYITCAPFNPAATESIA 476
        455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS 513
Qу
            Db
        477 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 536
        514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
Qу
           Db
        537 KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL 596
```

detect cancer in a patient and compositions comprising polypeptides,

CC

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574 CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVOPSVSPLEAPPPVSYDSIKLEPEN 633
Qу
           1:1:11
       597 CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIOPSSPLEA-SSVNYESIKHEPEN 655
Db
       634 PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVOETEAPYISIACDLIKETKLSTEPSPDF 692
Qу
           656 PPPYEEAMSVSLKKVSGIKEEIKEPENINAALOETEAPYISIACDLIKETKLSAEPAPDF 715
Db
       693 SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPOTOEEAVMLMKESLTEVS 752
Qу
           716 SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS 775
Db
       753 -ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF 809
Qу
           776 FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLOMEEL 835
Db
       810 NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS 868
Qу
           836 STAVYSNDDLFISKEAOIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS 895
Db
QУ
       869 DKSEIANIQSGADSLPCLELPCDLSFKNIYPK--DEVHVSDEFSENRSSVSKASISPSNV 926
                  896 HKSEIANAPDGAGSLPCTELPHDLSLKNIOPKVEEKISFSDDFSKNGSATSKVLLLPPDV 955
Db
       927 SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK 986
Qу
             956 SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK 1015
Db
Qу
       987 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIOAIOKSDEGHPFRA 1046
           1016 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA 1075
Db
Qу
       1047 YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1106
           1076 YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1135
Db
Qy
       1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
           Db
       1136 NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1192
RESULT 8
   AAY56967 standard; Protein; 1192 AA.
ХХ
AC
   AAY56967;
XX
DT
   25-APR-2000 (first entry)
XX
DE
   Human MAGI polypeptide.
XX
KW
   MAGI protein; neuroendocrine-specific protein; neuropathy; human;
KW
   spinal injury; neuronal degeneration; neuromuscular disorder; cancer;
KW
   psychiatric disorder; developmental disorder; inflammatory disorder;
```

stroke; cytostatic; cerebroprotective; neuroprotective.

KW XX

```
OS
    Homo sapiens.
XX
    WO200005364-A1.
PN
ХХ
    03-FEB-2000.
PD
XX
ΡF
    21-JUL-1999;
                  99WO-GB02360.
XX
PR
    22-JUL-1998;
                  98GB-0016024.
    19-JUL-1999:
                  99GB-0016898.
PR
XX
PA
    (SMIK ) SMITHKLINE BEECHAM PLC.
XX
PΙ
    Michalovich D, Prinjha RK;
XX
    WPI; 2000-182693/16.
DR
    N-PSDB; AAZ56886.
DR
XX
PT
    Novel polypeptides related to neuroendocrine-specific proteins and
    polynucleotides useful for diagnosis of various diseases and for
PT
    treatment of cancer and neurological disorders -
PT
XX
PS
    Claim 2; Page 20-21; 35pp; English.
XX
CC
    The invention relates to human MAGI protein, which is similar to
    neuroendocrine-specific protein. The MAGI protein can be expressed by
CC
    standard recombinant methodology. The MAGI polypeptides, polynucleotides
CC
    and antibodies are useful for treating diseases, including neuropathies,
CC
CC
    spinal injury, neuronal degeneration, neuromuscular disorders,
CC
    psychiatric disorders and developmental disorders, cancer, stroke and
CC
    inflammatory disorders. The polynucleoitde is also useful for chromosome
CC
    localization and for tissue expression studies. The present sequence
CC
    represents the human MAGI protein.
XX
SO
    Sequence 1192 AA;
                       75.2%; Score 4398.5; DB 21; Length 1192; 75.9%; Pred. No. 4.5e-221;
 Query Match
 Best Local Similarity
 Matches 908; Conservative 104; Mismatches 146; Indels
                                                          39; Gaps
                                                                     20;
Qy
           1 MEDIDOSSLVSSSTDSPPRPPPAFKYOFVTEPEDEEDEEEEEDEEDDEDLEELEVLERK 60
             Db
           1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDEDLEELEVLERK 58
          61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
Qу
             Dh
          59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPEROPSWDPSPVSSTVP 118
Qу
         116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
                  Db
         119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
Qу
         167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEOPGNTVSSGOEDFPSVLLETAASLPS 225
             179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEQPGNTISAGQEDFPSVLLETAASLPS 237
Db
         226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
QУ
```

Db	238	: :: :: : :: ::	297
Qу	286	SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVGKED	339
Db	298	SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNQQELPTALTKLVKED	357
Qу	340	RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANV	395
Db	358	EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL	416
Qy	396	ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT	454
Db	417	ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRPGAYITCAPFNPAATESIA	476
Qy	455	ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS	513
Db	477	TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT	536
Qу	514	KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL	573
Db	537	KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL	596
Qy	574	CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN	633
Db	597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN	655
Qy	634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF	692
Db	656	PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF	715
QУ	693	SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS	752
Db	716	,	775
Qy	753	-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF :: ::: :	809
Db	776	FESMI EYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKI PLQMEEL	835
Qy	810	NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS: :	868
Db	836	STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS	895
Qy	869	DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	926
Db	896	HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	955
Qу	927	SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	986
Db	956	SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK	1015
Qy	987	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1046
Db	1016	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1075
Qу	1047	YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1106

```
Db
         1076 YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1135
Qу
         1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
              Db
         1136 NGLTLLILALISLFSVPVIYERHOAOIDHYLGLANKNVKDAMAKIOAKIPGLKRKAE 1192
RESULT 9
AAB82349
     AAB82349 standard; Protein; 1192 AA.
ID
ХX
AC
     AAB82349;
XX
DT
     23-JUL-2001 (first entry)
XX
DE
     Human NOGO-A protein.
XX
KW
     NOGO-A; human; chromosome 2p21; neuropathy; spinal injury;
KW
     brain injury; stroke; neuronal degeneration; Alzheimer's disease:
     Parkinson's disease; neuromuscular disorder; psychiatric disorder;
KW
KW
     developmental disorder; neuroprotective; nootropic; neuroleptic;
     antiparkinsonian; cerebroprotective; neuroleptic; diagnosis;
KW
KW
     therapy.
XX
OS
     Homo sapiens.
XX
PN
     WO200136631-A1.
XX
PD
     25-MAY-2001.
XX
     14-NOV-2000; 2000WO-GB04345.
PF
XX
PR
     15-NOV-1999;
                   99GB-0026995.
PR
     24-JAN-2000; 2000GB-0001550.
XX
PA
     (SMIK ) SMITHKLINE BEECHAM PLC.
XX
PΙ
     Michalovich D, Prinjha R;
XX
DR
     WPI; 2001-343822/36.
DR
     N-PSDB: AAF90324.
XX
PΤ
     New polypeptide designated NOGO-C is a splice variant of the human NOGO
PT
     gene and may be useful in the treatment of neural disorders including
PΤ
     Alzheimer's and Parkinson's diseases -
XX
PS
     Disclosure; Page 26-27; 25pp; English.
XX
CC
     The present sequence is that of human NOGO-A. NOGO-A is a
CC
     previously known splice variant of the human NOGO gene on chromosome
CC
     2p21. The invention relates to a novel splice variant, NOGO-C (see
CC
     AAB82348). It provides NOGO-C polypeptides and polynucleotides, and
CC
     methods for producing such polypeptides by recombinant techniques.
CC
    Also disclosed are methods for utilising NOGO-C polypeptides and
CC
    polynucleotides in the treatment of diseases including neuropathies,
CC
     spinal injury, brain injury, stroke, neuronal degeneration, for
CC
     example Alzheimer's disease and Parkinson's disease, neuromuscular
```

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provided are methods for identifying agonists and agonists for
CC
CC
    use in treating conditions associated with NOGO-C imbalance, and
CC
    diagnostic assays for detecting diseases associated with
CC
    inappropriate NOGO-C activity or levels.
XX
SO
    Sequence
            1192 AA;
                    75.2%; Score 4398.5; DB 22; Length 1192;
 Query Match
 Best Local Similarity
                   75.9%; Pred. No. 4.5e-221;
 Matches 908; Conservative 104; Mismatches 146; Indels
                                                  39; Gaps
                                                           20;
         1 MEDIDQSSLVSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEDLEELEVLERK 60
Qу
           1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDLEELEVLERK 58
Db
        61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
Qу
           59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPEROPSWDPSPVSSTVP 118
Db
       116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
Qу
               119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPOAEPVWTPPAPAPAAPPSTPAAPKR 178
Db
Qу
       167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
           Db
       179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEQPGNTISAGQEDFPSVLLETAASLPS 237
Qу
       226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
           238 LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY 297
Db
Qу
       286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVG----KED 339
                 298 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNQQELPTALTKLVKED 357
Db
Qу
       340 RVVSPEKTMDIFNEMOMSVVAPVREEYADFKPFEOAWEVKDTYEGSRDVLAA----RANV 395
           358 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 416
Db
       396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
Qу
           Db
       417 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRPGAYITCAPFNPAATESIA 476
       455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS 513
Qу
           Dh
       477 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 536
Qу
       514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
           537 KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL 596
Db
       574 CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN 633
Qу
           Db
       597 CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIOPSSSPLEA-SSVNYESIKHEPEN 655
Qу
       634 PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF 692
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disorders, psychiatric disorders and developmental disorders. Also

CC

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656 PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF 715
Db
        693 SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS 752
QУ
           716 SDYSEMAKVEOPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPOKODETVMLVKESLTETS 775
Db
        753 -ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLOMEEF 809
QУ
            776 FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLOMEEL 835
Db
        810 NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS 868
Qу
           836 STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS 895
Db
        869 DKSEIANIQSGADSLPCLELPCDLSFKNIYPK--DEVHVSDEFSENRSSVSKASISPSNV 926
Qу
                 896 HKSEIANAPDGAGSLPCTELPHDLSLKNIOPKVEEKISFSDDFSKNGSATSKVLLLPPDV 955
Db
        927 SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK 986
Qу
           956 SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK 1015
Db
        987 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIOAIOKSDEGHPFRA 1046
Oy
           1016 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA 1075
Db
       1047 YLESEVAISEELVOKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1106
Qу
           1076 YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1135
Db
       1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
Qу
           Db
       1136 NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1192
RESULT 10
ABB81078
ID
    ABB81078 standard; Protein; 1192 AA.
XX
AC
    ABB81078;
XX
    05-NOV-2002 (first entry)
DT
XX
DE
    Human neurotransmitter receptor protein Nogo-A.
XX
KW
    Nerve regeneration; neuroprotection; neuronal degeneration; CNS; PNS;
KW
    central nervous system; peripheral nervous system; tranquillizer; Noqo;
KW
    vulnerary; cerebroprotective; anti-tumour; antidiabetic; anticonvulsant;
KW
    nootropic; antiparkinsonian; ophthalmological; analgesic; hepatotropic;
KW
    osteopathic; vasotropic; nephrotropic; cytostatic; antigen; gene therapy;
KW
    neurotransmitter receptor; human; receptor.
XX
OS
    Homo sapiens.
XX
PN
    US2002072493-A1.
```

XX

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PD
     13-JUN-2002.
XX
PF
     28-JUN-2001; 2001US-0893348.
XX
PR
     19-MAY-1998;
                    98IL-0124500.
PR
     21-JUL-1998:
                    98WO-US14715.
     22-DEC-1998:
                    98US-0218277.
PR
     19-MAY-1999; 99US-0314161.
XX
PA
     (YEDA ) YEDA RES & DEV CO LTD.
XX
ΡŤ
     Eisenbach-Schwartz M, Hauben E, Cohen IR, Beserman P, Mosonego A;
PΙ
     Moalem G;
XX
DR
     WPI; 2002-607255/65.
DR
     N-PSDB; ABN86601.
XX
     Promoting nerve regeneration and preventing neuronal degeneration in
PT
PT
     the central/peripheral nervous system from injury/disease, comprises
PT
     administering nervous system-specific activated T cells/antiqen, or
PΤ
     analogs/peptides
XX
PS
     Examples; Page 53-56; 93pp; English.
XX
CC
     The invention relates to promoting nerve regeneration or conferring
CC
     neuroprotection and preventing or inhibiting neuronal degeneration in the
CC
     central/peripheral nervous system (NS). The method involves administering
CC
     NS-specific activated T cells, NS-specific antigen, its analogue or its
CC
     peptide, a nucleotide sequence the NS-specific antigen or its analogue or
CC
     combinations. The method is useful for promoting nerve regeneration and
CC
     preventing neuronal degeneration in central/peripheral nervous system
CC
     from injury/disease, where the injury is spinal cord injury, blunt
CC
     trauma, penetrating trauma, hemorrhagic stroke, ischaemic stroke or
CC
     damages caused by surgery such as tumour excision. The disease is not an
CC
     autoimmune disease or neoplasm. The disease results in a degenerative
CC
     process occurring in either gray or white matter or both. The disease
CC
     is diabetic neuropathy, senile dementia, Alzheimer's disease, Parkinson's
CC
     disease, facial nerve (Bell's) palsy, glaucoma, Huntington's chorea,
CC
     amyotrophic lateral sclerosis, non-arteritic optic neuropathy, and
CC
     vitamin deficiency, intervertebral disc herniation, prion diseases such
CC
     as Creutzfeldt-Jakob disease, carpal tunnel syndrome, peripheral
CC
     neuropathies associated with various diseases, including but not limited
CC
     to uremia, porphyria, hypoglycemia, Sjorgren Larsson syndrome, acute
CC
     sensory neuropathy, chronic ataxic neuropathy, biliary cirrhosis, primary
CC
     amyloidosis, obstructive lung diseases, acromegaly, malabsorption
CC
     syndromes, polycythemia vera, immunoglobulin (Ig)A- and IgG gamma-
CC
     pathies, complications of various drugs (e.g., metronidazole) and toxins
CC
     (e.g., alcohol or organophosphates), Charcot-Marie-Tooth disease, ataxia
CC
     telangectasia, Friedreich's ataxia, amyloid polyneuropathies,
CC
     adrenomyeloneuropathy, Giant axonal neuropathy, Refsum's disease, Fabry's
CC
     disease, or lipoproteinemia. The present sequence represents the human
CC
     neurotransmitter receptor protein Nogo-A, an example of NS-specific
CC
     antigen.
XX
SQ
     Sequence
               1192 AA;
```

	Best Loc Matches		Similarity 75.9%; Pred. No. 4.5e-221; 8; Conservative 104; Mismatches 146;	Indels	39;	Gaps	20
Qу		1	MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEE				60
Db		1	:	: :: EEEEEDEDEI	DLEEL	 EVLERK	58
Qу		61	PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPI				115
Db		59					118
Qу		116	APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE				166
Db		119				 PAAPKR	178
Qу		167	RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVS				225
DŁ		179	RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEQPGNTIS				237
Qу		226	LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPI				285
Db		238	LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVS				297
Qу		286	SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSA			KED	339
DŁ		298	SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSN			KLVKED	357
Qу		340	RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKI			RANV : :	395
Dk		358	EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKI				416
Qу		396	ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDS	SSRAYITCA:		ATESTT	454
Db		417	ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKD				476
Qу		455	ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNI				513
Db		477	TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNI				536
Qу		514	KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKV				573
Db		537	KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKI				596
Qу		574	CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVS				633
Db		597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSS				655
Qу		634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYIS				692
Db		656	PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYIS				715
Qу		693	SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEV				752
Db			SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDV	VPQKQDETVI	MLVKE	SLTETS	775
Qу		753	-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-AS				809
			,				

```
Db
        776 FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLQMEEL 835
        810 NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS 868
Qу
            836 STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS 895
Db
        869 DKSEIANIQSGADSLPCLELPCDLSFKNIYPK--DEVHVSDEFSENRSSVSKASISPSNV 926
Qу
                    896 HKSEIANAPDGAGSLPCTELPHDLSLKNIOPKVEEKISFSDDFSKNGSATSKVLLLPPDV 955
Db
        927 SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK 986
Qу
               Db
        956 SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK 1015
        987 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA 1046
Qу
            Db
       1016 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA 1075
       1047 YLESEVAISEELVOKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1106
Qу
            1076 YLESEVAISEELVOKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1135
Db
Qу
       1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
            Db
       1136 NGLTLLILALISLFSVPVIYERHOAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1192
RESULT 11
ABG30938
    ABG30938 standard; Protein; 1192 AA.
XX
AC
    ABG30938;
XΧ
DT
    21-OCT-2002 (first entry)
XX
DE
    Human NogoA protein.
XX
KW
    Human; Nogo; BACE; acute neuronal injury; spinal injury; head injury;
KW
    stroke; peripheral nerve damage; neoplastic disorder; glioblastoma;
KW
    neuroblastoma; hyperproliferative disorder; dysproliferative disorder;
KW
    cirrhosis; psoriasis; keloid formation; fibrocystic condition; cancer;
    tissue hypertrophy; central nervous system; axon regeneration; NogoA;
KW
KW
    Nogo-associated disease; metastasis.
XX
OS
    Homo sapiens.
XX
ÞΝ
    WO200257483-A2.
ХX
PD
    25-JUL-2002.
XX
PF
    18-JAN-2002; 2002WO-GB00228.
XX
PR
    18-JAN-2001; 2001GB-0001312.
XX
PΑ
    (GLAX ) GLAXO GROUP LTD.
PΑ
    (SMIK ) SMITHKLINE BEECHAM PLC.
XX
```

```
Blackstock WP, Hale RS, Prinjha R, Rowley A;
PΤ
XX
DR
    WPI; 2002-599722/64.
    N-PSDB; ABK90134.
DR
XX
PT
    Identifying modulators of Nogo or BACE activity for treating acute
PT
    neuronal injuries, neoplastic or dysproliferative disorders, comprises
PT
    providing and monitoring interaction between Nogo and BACE polypeptides
PT
XX
PS
    Disclosure; Page 59-62; 68pp; English.
XX
CC
    The present invention relates to a new method of identifying modulators
CC
    of Nogo function or BACE activity. The method involves providing Nogo and
CC
    BACE polypeptides capable of binding with each other, monitoring the
CC
    interaction between these polypeptides, and determining if the test agent
CC
    is a modulator of Nogo or BACE activity. The method is useful in treating
CC
    acute neuronal injuries, such as spinal or head injury, stroke,
CC
    peripheral nerve damage, and in neoplastic (e.g. glioblastomas,
CC
    neuroblastomas), hyperproliferative or dysproliferative disorders (e.g.
CC
    cirrhosis, psoriasis, keloid formation, fibrocystic conditions, tissue
CC
    hypertrophy) of the central nervous system. The BACE polypeptide is
CC
    useful in screening methods to identify agents that may act as modulators
CC
    of BACE activity and in particular agents that may be useful in treating
CC
    Nogo-associated diseases. The modulators of Nogo or BACE polypeptides,
CC
    and the polynucleotide encoding the BACE polypeptide are useful in
CC
    manufacturing a medicament for the treatment or prevention of disorders
    responsive to the modulation of Nogo activity, in alleviating the
CC
    symptoms or improving the condition of a patient suffering from this
CC
CC
    disorder, in axon regeneration, or in preventing metastasis or spreading
CC
    of a cancer. The polynucleotide may also be an essential component in
CC
    assays, a probe, in recombinant protein synthesis, and in gene therapy
CC
    techniques. The present amino acid sequence represents the human NogoA
CC
    protein of the invention.
XX
SO
    Sequence
              1192 AA;
  Query Match
                        75.2%; Score 4398.5; DB 23; Length 1192;
  Best Local Similarity
                       75.9%; Pred. No. 4.5e-221;
 Matches 908; Conservative 104; Mismatches 146; Indels
           1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
Qy
             Db
           1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDDLEELEVLERK 58
Qу
          61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
             59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP 118
Db
Qу
         116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
                  119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
Db
         167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Qу
             Db
         179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEQPGNTISAGQEDFPSVLLETAASLPS 237
```

Qy Db		LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY : : : : :	
Qy	286	SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVGKED	339
Db	298	:: : : : : : :	357
Qу	340	RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANV	395
Db	358	::: :	416
Qy	396	ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT	454
Db	417	: : : :	476
Qy	455	ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS	513
Db	477	TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT	536
Qу	514	KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL	573
Db	537	KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL	596
Qу	574	CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN	633
Db	597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN	655
Qy	634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF	692
Db		PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF	
Qy	693	SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS : : : : : : : :	752
Db		SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS	
Qу		-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF	
Db		FESMI EYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKI PLQMEEL	
Qу		NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS : : : :	
Db		STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS	
Qу		DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	
Db		HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	
Qу		SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	
Db		SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK	
Qy		KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	
Db Ov		KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA VLESEVAISEELVOKYSNSALGHVNSTIKELPRIELVDDLVDSLKRAVLMWVFTVVGALR	
LIV	104/	- 1 DB-5B-VA L5B-B-LOUDK Y 5N/5A-D-B-WN/5-L-K-B-D-R-D-B-L-WDHLA/DS-L-K-B-A-VL-WW/7-B-T-V-WC-A-L-B-	1 1 1 1 1 1 1 1 1 1 1 1

```
Db
        1076 YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1135
        1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
QУ
             Dh
        1136 NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1192
RESULT 12
AAY71311
     AAY71311 standard; Protein; 1178 AA.
ID
ХX
AC
    AAY71311;
XX
DT
     02-NOV-2000 (first entry)
XX
DE
     Human neurite growth inhibitor Nogo.
XX
KW
     Human; neurite growth inhibitor; Nogo; neural cell; myelin; CNS;
KW
     central nervous system; neoplastic disease; antiproliferative; glioma;
     antisense gene therapy; neuroblastoma; menagioma; retinoblastoma;
KW
KW
     degenerative nerve disease; Alzheimer's disease; Parkinson's disease;
KW
     hyperproliferative disorder; beniqn dysproliferative disorder; diagnosis;
     psoriasis; tissue hypertrophy; neuronal regeneration; treatment;
KW
KW
     structural plasticity; screening.
XX
OS
     Homo sapiens.
XX
FΗ
     Key
                    Location/Qualifiers
FT
     Misc-difference 187
FT
                    /label= Unknown
FT
    Misc-difference 188
FT
                    /label= Unknown
FT
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FT
                    /label= Unknown
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FT
    Misc-difference 477
FT
                    /label= Unknown
FT
    Region
                    994..1174
FT
                    /note= "Region specifically described in claim 16"
FT
    Region
                    977..1012
FT
                    /note= "Region specifically described in claim 16"
FT
    Region
                    1079..1114
FT
                    /note= "Region specifically described in claim 16"
XX
ΡN
    WO200031235-A2.
ХX
PD
    02-JUN-2000.
XX
PF
    05-NOV-1999;
                   99WO-US26160.
XX
PR
```

06-NOV-1998;

98US-0107446.

```
XX
     (SCHW/) SCHWAB M E.
PA
     (CHEN/) CHEN M S.
PΑ
XX
ΡI
    Schwab ME,
                Chen MS;
XX
    WPI: 2000-400052/34.
DR
XX
PT
    Nogo proteins and nucleic acids useful for treating neoplastic
PT
    disorders of the central nervous system and inducing regeneration of
PT
    neurons -
XX
PS
    Claim 11; Fig 13; 122pp; English.
XX
CC
    The present sequence is a human Nogo protein which is a
CC
    potent neural cell growth inhibitor and is free of all central nervous
CC
    system (CNS) myelin material with which it is natively associated. The
CC
    human Nogo sequence was derived by aligning human expressed sequence tags
CC
     (ESTs) e.g. AA158636, AA333267, AA081783, AA167765, AA322918, AA092565,
CC
    AA081525 and AA081840 with the rat Nogo sequence.
CC
    Nogo proteins and fragments displaying neurite growth inhibitory
CC
    activity are used in the treatment of neoplastic disease of the CNS
CC
    e.g. glioma, glioblastoma, medulloblastoma, craniopharyngioma, ependyoma,
CC
    pinealoma, haemangioblastoma, acoustic neuroma, oligodendroglioma,
CC
    menagioma, neuroblastoma or retinoblastoma and degenerative nerve
CC
    diseases e.q. Alzheimer's and Parkinson's diseases. Therapeutics which
CC
    promote Nogo activity can be used to treat or prevent hyperproliferative
CC
    or benign dysproliferative disorders e.g. psoriasis and tissue
CC
    hypertrophy. Ribozymes or antisense Nogo nucleic acids can be used to
CC
    inhibit production of Nogo protein to induce regeneration of neurons or
CC
    to promote structural plasticity of the CNS in disorders where neurite
    growth, regeneration or maintenance are deficient or desired.
CC
    The animal models can be used in diagnostic and screening methods for
CC
CC
    predisposition to disorders and to screen for or test molecules which
CC
    can treat or prevent disorders or diseases of the CNS.
CC
    Note: SEQ ID numbers 35-42 are referred in claim 32 and SEQ ID NO: 29
CC
    in disclosure of the specification. However the specification does not
CC
    include sequences for these SEQ ID numbers.
XX
SO
    Sequence
               1178 AA;
 Query Match
                        73.1%; Score 4276.5; DB 21; Length 1178;
  Best Local Similarity
                        73.8%; Pred. No. 1e-214;
 Matches 883; Conservative 104; Mismatches 158; Indels
                                                            51; Gaps
Qу
           1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
             1 MEDLDQSPLVSSS-DSVPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDLEELEVLERK 58
Db
Qу
          61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
             59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPEROPSWDPSPVSSTVP 118
Db
Qу
         116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
                   Db
         119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
```

Qy Db		RGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSL	
Qу	227	SPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYS	286
Db	225	SPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEYS	284
Qy	287	EMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVGKEDR	340
Db	285	EMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHXQQELPTALTKLVKEDE	344
Qy	341	VVSPEKTMDI FNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVE ::: :	396
Db	345	VVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNLE	403
Qу	397	SKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTTA	455
Db	404	SKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRSGAYITCAPFNPAATESIAT	463
Qy	456	NTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLSK	514
Db	464	NIFPLLEDPTSENXTDEKKIEEKKAQIVTEKNTSTKTSNPFFVAAQDSCTDYVTTDNLTK	523
Qy	515	VTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLC	574
Db	524	VTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQLC	583
Qy	575	PSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENP	634
Db	584	PSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPENP	642
Qу		PPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSN	
Db		PPYEEAMSVSLKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDFSD	
Qу		YSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS-E	
Db		YSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETSFE	
Qу		TVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEFNT::::: :	
Db	763	SMIEYENKEKLSALPPEGGKPYLESFKLSLVNTKDTLLPDEVSTLSKKEKIPLQMEELST	822
Qу		AIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVSDK :	
Db		AVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVSHK	
Qу	871	SEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSA	928
Db		SEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDVSA	
Qу		L-EPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKK	
Db		LGHTQAEIESIVKPKVLEKEAEKKLPSDTEKEDRSPSAIFSADLGKTSVVDLLYWRDIKK	
Qу	988	${\tt TGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAY}$	1047

```
1003 TGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAY 1062
Db
        1048 LESEVAISEELVOKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFN 1107
Qу
             1063 LESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFN 1122
Db
Qу
        1108 GLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
             1123 GLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1178
Dh
RESULT 13
AAU33228
    AAU33228 standard; Protein; 1246 AA.
ХX
AC
    AAU33228;
XX
DТ
    18-DEC-2001 (first entry)
XX
    Novel human secreted protein #3719.
DE
XX
KW
    Human; vaccination; gene therapy; nutritional supplement;
     stem cell proliferation; haematopoiesis; nerve tissue regeneration;
KW
KW
     immune suppression; immune stimulation; anti-inflammatory; leukaemia.
XX
OS
    Homo sapiens.
XX
PN
    WO200179449-A2.
XX
PD
    25-OCT-2001.
XX
PF
    16-APR-2001; 2001WO-US08656.
XX
PR
    18-APR-2000; 2000US-0552929.
PR
    26-JAN-2001; 2001US-0770160.
XX
     (HYSE-) HYSEQ INC.
PΑ
XX
PΙ
    Tang YT, Liu C, Drmanac RT;
XX
DR
    WPI; 2001-611725/70.
XX
PT
    Nucleic acids encoding a range of human polypeptides, useful in genetic
PT
    vaccination, testing and therapy -
XX
PS
    Claim 20; Page 737; 765pp; English.
XX
    The invention relates to novel human secreted polypeptides. The
CC
CC
    polypeptides and antibodies to the polypeptides are useful for
CC
    determining the presence of or predisposition to a disease associated
CC
    with altered levels of polypeptide. The polypeptides are also useful for
CC
    identifying agents (agonists and antagonists) that bind to them. Cells
CC
    expressing the proteins are useful for identifying a therapeutic agent
CC
    for use in treatment of a pathology related to aberrant expression or
CC
    physiological interactions of the polypeptide. Vectors comprising
CC
    the nucleic acids encoding the polypeptides and cells genetically
```

```
CC
    The proteins are useful in genetic vaccination, testing and
CC
    therapy, and can be used as nutritional supplements. They may be used to
CC
    increase stem cell proliferation; to regulate haematopoiesis; and in
CC
    bone, cartilage, tendon and/or nerve tissue growth or regeneration;
CC
    immune suppression and/or stimulation; as anti-inflammatory agents; and
    in treatment of leukaemias. AAU29510-AAU33304 represent the amino acid
CC
CC
    sequences of novel human secreted proteins of the invention.
XX
            1246 AA;
SO
    Sequence
 Query Match
                    70.4%; Score 4116; DB 22;
                                           Length 1246;
 Best Local Similarity
                    72.0%; Pred. No. 2.6e-206;
 Matches 873; Conservative 112; Mismatches 171;
                                           Indels
                                                   56; Gaps
                                                            27;
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
Qу
           Db
         42 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDEDLEELEVLERK 99
         61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPEROPSWERSPAA---P 115
Qу
           Db
        100 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPEROPSWDPSPVSSTVP 159
        116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
Qу
                Db
        160 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 219
        167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Qу
           Db
        220 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEOPGNTISAGOEDFPSVLLETAASLPS 278
        226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
Qу
           279 LSPLSAASFKEHEYLGNLSTVLPTEGTLOENVSEASKEVSEKAKTLLIDRDLTEFSELEY 338
Db
Qу
        286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVG----KED 339
           Db
        339 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNOOELPTALTKLVKED 398
        340 RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAA----RANV 395
Qу
            399 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 457
Db
        396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
Qу
           458 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRSGAYITCAPFNPAATESIA 517
Db
Qy
       455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVODSEADYVTTDTLS 513
            Db
       518 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 577
Qу
       514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
               Db
       578 KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL 637
       574 CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN 633
Qу
```

engineered to express them are also useful for producing the proteins.

CC

```
638 CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN 696
Db
        634 PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF 692
Qу
           697 PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF 756
Db
        693 SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS 752
Qу
           757 SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS 816
Db
        753 -ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLOMEEF 809
Qу
            817 FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLQMEEL 876
Db
        810 NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS 868
Qу
           877 STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS 936
Db
        869 DKSEIANIQSGADSLPCLELPCDLSFKNIYPK--DEVHVSDEFSENRSSVSKASISPSNV 926
Qу
                   937 HKSEIANAPDGAGSLPCTELPHDLSLKNIOPKVEEKISFSDDFSKNGSATSKVLLLPPDV 996
Db
        927 SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK 986
Qу
              997 SALATOAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK 1056
Db
Qу
        987 KTGVVFGAS-LFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFR 1045
           1057 KTGVVFGASAVFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFR 1116
Db
       1046 AY---LESEVAISEELVQKYSNSALGHV-NSTIKELRR---LFLVDDLVDSLK-FAVLMW 1097
Qу
               Db
       1117 AISGNLESCLYLRELGSGRYSNSALGSMWNCTVKGNFRAPSFFSWMDLVDSLRSFAVLMW 1176
Qу
       1098 VFTYVGALFNGLTLL-----ILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKI 1151
           ||||||
                            1177 VFTYVGCL--GLMVLDTTGFWALNFISSSGSWLIYERHQAQIDHYLGLANKNVKDAMAKI 1234
Db
Qу
       1152 OAKIPGLKRKAD 1163
           1111111:
Db
       1235 QAKIPGLKRKAE 1246
RESULT 14
AAY71562
ID
   AAY71562 standard; Protein; 803 AA.
XX
AC
   AAY71562;
XX
DT
    02-NOV-2000 (first entry)
XX
DΕ
   Rat Nogo A protein fragment used in the construction of mutant NiG.
XX
KW
   Rat; neurite growth inhibitor; Nogo A; neural cell; myelin; CNS;
KW
   central nervous system; neoplastic disease; antiproliferative; glioma;
KW
   antisense gene therapy; neuroblastoma; menagioma; retinoblastoma;
```

degenerative nerve disease; Alzheimer's disease; Parkinson's disease;

KW

KW hyperproliferative disorder; benign dysproliferative disorder; diagnosis; KW psoriasis; tissue hypertrophy; neuronal regeneration; treatment; structural plasticity; screening; mutant; mutein. KW XX OS Rattus sp. XX PNWO200031235-A2. XX PD02-JUN-2000. XX PF 05-NOV-1999; 99WO-US26160. XX PR 06-NOV-1998; 98US-0107446. XX PA(SCHW/) SCHWAB M E. PA(CHEN/) CHEN M S. XXPΙ Schwab ME, Chen MS: XX DR WPI; 2000-400052/34. XX Nogo proteins and nucleic acids useful for treating neoplastic PTPΤ disorders of the central nervous system and inducing regeneration of PTneurons -XX PS Example; Page -; 122pp; English. XX The patent relates to neurite growth inhibitor Nogo which is free of CCCC all central nervous system (CNS) myelin material with which it is CC natively associated. Nogo proteins and fragments displaying neurite CC growth inhibitory activity are used in the treatment of neoplastic CC disease of the CNS e.g. glioma, glioblastoma, medulloblastoma, CC craniopharyngioma, ependyoma, pinealoma, haemangioblastoma, acoustic CC neuroma, oligodendroglioma, menagioma, neuroblastoma or retinoblastoma CC and degenerative nerve diseases e.g. Alzheimer's and Parkinson's CC diseases. Therapeutics which promote Nogo activity can be used to treat CC or prevent hyperproliferative or benign dysproliferative disorders e.g. psoriasis and tissue hypertrophy. Ribozymes or antisense Nogo nucleic CC CC acids can be used to inhibit production of Nogo protein to induce CC regeneration of neurons or to promote structural plasticity of the CNS CC in disorders where neurite growth, regeneration or maintenance are CCdeficient or desired. The animal models can be used in diagnostic and CC screening methods for predisposition to disorders and to screen for or CC test molecules which can treat or prevent disorders or diseases of the CC CNS. The present sequence is a fragment of rat Nogo A protein shown in

CC Nogo protein. Major inhibitory region was identified in the CC Nogo A sequence from amino acids 172-974, particularly amino acids CC542-722. In addition, N-terminal region 1-171 was found to be inhibitory CC to NIH 3T3 fibroblast spreading. CC

is composed of His-tag/T7-tag/Nogo-A sequence aa 172-974/His-tag.

AAY71310, which is used in the construction of mutant NiG. The mutant

Nogo A deletion mutants were used for mapping the inhibitory sites of

CC

CC

CC

Note: The present sequence is not given in the specification but is CC derived from rat Nogo A sequence shown in AAY71310. SEQ ID numbers 35-42 CC are referred in claim 32 and SEQ ID NO: 29 in disclosure of the specification. However, the specification does not include sequences for

CC CCthese SEO ID numbers.

Query Match 68.8%; Score 4023; DB 21; Length 803; 99.9%; Best Local Similarity Pred. No. 1.1e-201; Matches 802; Conservative Mismatches 0; Indels Gaps 0; 0; Qу 172 VDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLST 231 Db 1 VDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLST 60 Qу 232 VSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSS 291 Db VSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSS 120 292 FKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIF 351 Qу 121 FKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIF 180 Db Qу 352 NEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEOK 411 181 NEMOMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCIEDSLEQK 240 Db Qу 412 SLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTD 471 Db 241 SLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTD 300 Qу 472 EKKIEERKAQİITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDL 531 Db 301 EKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDL 360 Qу 532 VQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDI 591 361 VQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDI 420 Db Qу 592 VMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTK 651 Db 421 VMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTK 480 Qy 652 EGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAEL 711 Db 481 EGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAEL 540 712 VEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELG 771 Qу Db 541 VEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELG 600 772 KPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESE 831 Qу 601 KPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESE 660 Db 832 TFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCD 891 Qу Db 661 TFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCD 720 892 LSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKK 951 Qу

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Dh
RESULT 15
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XX
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     AAY71386;
XX
DT
     02-NOV-2000 (first entry)
XX
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     Rat Nogo A protein fragment used in the construction of mutant NiG-D1.
XX
KW
     Rat; neurite growth inhibitor; Nogo A; neural cell; myelin; CNS;
KW
     central nervous system; neoplastic disease; antiproliferative; glioma;
KW
     antisense gene therapy; neuroblastoma; menagioma; retinoblastoma;
KW
     degenerative nerve disease; Alzheimer's disease; Parkinson's disease;
KW
     hyperproliferative disorder; benign dysproliferative disorder; diagnosis;
KW
     psoriasis; tissue hypertrophy; neuronal regeneration; treatment;
KW
     structural plasticity; screening; mutant; mutein.
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PD
     02-JUN-2000.
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PF
     05-NOV-1999;
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PR
     06-NOV-1998;
                   98US-0107446.
XX
PΑ
     (SCHW/) SCHWAB M E.
     (CHEN/) CHEN M S.
PΑ
XX
PΙ
     Schwab ME, Chen MS;
XX
DR
     WPI; 2000-400052/34.
XX
PT
     Nogo proteins and nucleic acids useful for treating neoplastic
PT
     disorders of the central nervous system and inducing regeneration of
PT
     neurons -
XX
PS
     Example; Page -; 122pp; English.
XX
CC
     The patent relates to neurite growth inhibitor Nogo which is free of
CC
     all central nervous system (CNS) myelin material with which it is
CC
     natively associated. Nogo proteins and fragments displaying neurite
CC
     growth inhibitory activity are used in the treatment of neoplastic
CC
     disease of the CNS e.g. glioma, glioblastoma, medulloblastoma,
CC
     craniopharyngioma, ependyoma, pinealoma, haemangioblastoma, acoustic
CC
     neuroma, oligodendroglioma, menagioma, neuroblastoma or retinoblastoma
CC
     and degenerative nerve diseases e.g. Alzheimer's and Parkinson's
CC
     diseases. Therapeutics which promote Noqo activity can be used to treat
```

CC or prevent hyperproliferative or benign dysproliferative disorders e.g. CC psoriasis and tissue hypertrophy. Ribozymes or antisense Nogo nucleic acids can be used to inhibit production of Nogo protein to induce CCregeneration of neurons or to promote structural plasticity of the CNS CC CCin disorders where neurite growth, reqeneration or maintenance are deficient or desired. The animal models can be used in diagnostic and CCscreening methods for predisposition to disorders and to screen for or CC CC test molecules which can treat or prevent disorders or diseases of the CC CNS. The present sequence is a fragment of rat Nogo A protein shown in CCAAY71310, which is used in the construction of mutant NiG-D1. NiG-D1 CC is composed of His-tag/T7-tag/Nogo-A sequence aa 172-908/vector. CC Nogo A deletion mutants were used for mapping the inhibitory sites of CC Nogo protein. Major inhibitory region was identified in the CCNogo A sequence from amino acids 172-974, particularly amino acids CC542-722. In addition, N-terminal region 1-171 was found to be inhibitory CCto NIH 3T3 fibroblast spreading. CCNote: The present sequence is not given in the specification but is CC derived from rat Nogo A sequence shown in AAY71310. SEQ ID numbers 35-42 CCare referred in claim 32 and SEQ ID NO: 29 in disclosure of the CCspecification. However, the specification does not include sequences for CCthese SEQ ID numbers. XX

63.5%;

SQ Sequence 737 AA;

Query Match

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Score 3714; DB 21;
                                         Length 737;
 Best Local Similarity
                   99.9%;
                         Pred. No. 1.2e-185;
 Matches 736; Conservative
                        1; Mismatches
                                      0;
                                         Indels
                                                 0; Gaps
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           Db
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Qу
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Qу
          Db
       121 FKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIF 180
Qу
       352 NEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQK 411
           Db
       181 NEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCIEDSLEOK 240
       412 SLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTD 471
Qу
          241 SLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTD 300
Db
Qу
       472 EKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDL 531
          Db
       301 EKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDL 360
Qу
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          Db
       361 VQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDI 420
       592 VMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTK 651
Qу
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Db	421	
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Db	721	LSFKNIYPKDEVHVSDE 737

Search completed: December 19, 2003, 15:35:11 Job time : 52 secs

GenCore version 5.1.6 Copyright (c) 1993 - 2003 Compugen Ltd.

OM protein - protein search, using sw model

Run on:

December 19, 2003, 15:34:13 ; Search time 21 Seconds

(without alignments)

2343.216 Million cell updates/sec

Title:

US-09-830-972-2

Perfect score:

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Sequence:

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Scoring table:

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Gapop 10.0 , Gapext 0.5

Searched:

328717 seqs, 42310858 residues

Total number of hits satisfying chosen parameters:

328717

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

Issued Patents AA:*

- 1: /cgn2 6/ptodata/1/iaa/5A COMB.pep:*
- 2: /cgn2 6/ptodata/1/iaa/5B COMB.pep:*
- 3: /cgn2 6/ptodata/1/iaa/6A COMB.pep:*
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- 6: /cgn2 6/ptodata/1/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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Result		Query				
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1	908	15.5	199	2	US-08-700-607-1	Sequence 1, Appli
2	789.5	13.5	776	2	US-08-700-607-5	Sequence 5, Appli
3	716.5	12.3	356	2	US-08-700-607-6	Sequence 6, Appli
4	685	11.7	208	2	US-08-700-607 - 7	Sequence 7, Appli
5	671	11.5	267	2	US-08-700-607-8	Sequence 8, Appli
6	539.5	9.2	168	4	US-09-149-476-563	Sequence 563, App
7	519	8.9	241	2	US-08-700-607-3	Sequence 3, Appli
8	302.5	5.2	8991	4	US-08-714-741-32	Sequence 32, Appl
9	286	4.9	92	4	US-09-149-476-411	Sequence 411, App
10	279.5	4.8	1786	3	US-08-973-462-8	Sequence 8, Appli
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14	254.5	$\frac{4.3}{4.4}$	1596	4		Sequence 4463, Ap
15	234.3			2	US-08-978-277A-4	Sequence 4, Appli
16	233	4.0	1142		US-08-993-118-7	Sequence 7, Appli
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17	233	4.0	1142	4	US-09-066-281B-7	Sequence 7, Appli
18	232.5	4.0	1805	1	US-07-853-913-2	Sequence 2, Appli
19	228	3.9	1142	3	US-09-061-709-2	Sequence 2, Appli
20	228	3.9	1142	4	US-09-899-651-2	Sequence 2, Appli
21	227	3.9	1270	4	US-07-757-022B-44	Sequence 44, Appl
22	227	3.9	1311	4	US-07-757-022B-42	Sequence 42, Appl
23	227	3.9	1313	4	US-07-757-022B-142	Sequence 142, App
24	227	3.9	1314	4	US-07-757-022B-50	Sequence 50, Appl
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26	227	3.9	1320	4	US-07-757-022B-60	Sequence 60, Appl
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28	227	3.9	1361	4	US-07-757-022B-40	Sequence 40, Appl
29	227	3.9	1363	4	US-07-757-022B-52	Sequence 52, Appl
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31	227	3.9	1404	4	US-07-757-022B-62	Sequence 62, Appl
32	225	3.8	688	3	US-09-141-047-8	Sequence 8, Appli
33	224.5	3.8	941	4	US-07-757-022B-14	Sequence 14, Appl
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36	224.5	3.8	1049	4	US-07-757-022B-58	Sequence 58, Appl
37	224.5	3.8	1140	4	US-07-757-022B-104	Sequence 104, App
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39	224.5	3.8	1346	4	US-08-978-277A-2	Sequence 2, Appli
40	224	3.8	630	3	US-08-973-462-9	Sequence 9, Appli
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45	218.5	3.7	2842	1	US-07-741-940-7	Sequence 40, Appl
10	210. 5	J. 1	2042	_	03-0/-/41-340-/	Sequence 7, Appli

ALIGNMENTS

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RESULT 1
US-08-700-607-1
; Sequence 1, Application US/08700607
; Patent No. 5858708
; GENERAL INFORMATION:
    APPLICANT: Bandman, Olga
    APPLICANT: Au-Young, Janice
    APPLICANT: Goli, Surya K.
    APPLICANT: Hillman, Jennifer L.
    TITLE OF INVENTION: TWO NOVEL HUMAN NSP-LIKE PROTEINS
    NUMBER OF SEQUENCES: 9
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: U.S.
      ZIP: 94304
```

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COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ Version 1.5
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/700,607
      FILING DATE: Filed Herewith
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0114 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
   INFORMATION FOR SEO ID NO: 1:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 199 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
    IMMEDIATE SOURCE:
      LIBRARY:
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US-08-700-607-1
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US-08-700-607-5
; Sequence 5, Application US/08700607
; Patent No. 5858708
  GENERAL INFORMATION:
    APPLICANT: Bandman, Olga
    APPLICANT: Au-Young, Janice
    APPLICANT: Goli, Surya K.
    APPLICANT: Hillman, Jennifer L.
```

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TITLE OF INVENTION: TWO NOVEL HUMAN NSP-LIKE PROTEINS
     NUMBER OF SEQUENCES: 9
     CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: U.S.
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ Version 1.5
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/700,607
      FILING DATE: Filed Herewith
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0114 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
   INFORMATION FOR SEQ ID NO: 5:
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      CLONE: 307307
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US-08-700-607-6
; Sequence 6, Application US/08700607
 Patent No. 5858708
  GENERAL INFORMATION:
    APPLICANT: Bandman, Olga
    APPLICANT: Au-Young, Janice
    APPLICANT: Goli, Surya K.
    APPLICANT: Hillman, Jennifer L.
    TITLE OF INVENTION: TWO NOVEL HUMAN NSP-LIKE PROTEINS
    NUMBER OF SEQUENCES: 9
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Incyte Pharmaceuticals, Inc.
     STREET: 3174 Porter Drive
     CITY: Palo Alto
     STATE: CA
     COUNTRY: U.S.
     ZIP: 94304
    COMPUTER READABLE FORM:
     MEDIUM TYPE: Diskette
     COMPUTER: IBM Compatible
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OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ Version 1.5
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/700,607
     FILING DATE: Filed Herewith
    ATTORNEY/AGENT INFORMATION:
     NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
     REFERENCE/DOCKET NUMBER: PF-0114 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO: 6:
    SEQUENCE CHARACTERISTICS:
     LENGTH: 356 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
    IMMEDIATE SOURCE:
     LIBRARY: GenBank
      CLONE: 307309
US-08-700-607-6
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        883 LPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPOTEMGSIVKSK 942
QУ
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         93 RRGLAEPG--SFLD-YPSTEPQPGPEL------PPGDGALEPETPM----- 129
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RESULT 4

US-08-700-607-7

[;] Sequence 7, Application US/08700607

[;] Patent No. 5858708

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GENERAL INFORMATION:
     APPLICANT: Bandman, Olga
     APPLICANT: Au-Young, Janice
     APPLICANT: Goli, Surya K.
     APPLICANT: Hillman, Jennifer L.
     TITLE OF INVENTION: TWO NOVEL HUMAN NSP-LIKE PROTEINS
     NUMBER OF SEQUENCES: 9
     CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: U.S.
      ZIP: 94304
     COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ Version 1.5
     CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/700,607
      FILING DATE: Filed Herewith
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0114 US
     TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
   INFORMATION FOR SEQ ID NO: 7:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 208 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
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      CLONE: 307311
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Qу
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; Sequence 8, Application US/08700607
; Patent No. 5858708
  GENERAL INFORMATION:
    APPLICANT: Bandman, Olga
    APPLICANT: Au-Young, Janice
    APPLICANT: Goli, Surya K.
    APPLICANT: Hillman, Jennifer L.
    TITLE OF INVENTION: TWO NOVEL HUMAN NSP-LIKE PROTEINS
    NUMBER OF SEQUENCES: 9
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: U.S.
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEO Version 1.5
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/700,607
      FILING DATE: Filed Herewith
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0114 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO: 8:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 267 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
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      CLONE: 281046
US-08-700-607-8
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; Sequence 563, Application US/09149476
; Patent No. 6420526
; GENERAL INFORMATION:
  APPLICANT: Rosen et al.
  TITLE OF INVENTION: 186 Human Secreted proteins
  FILE REFERENCE: PZ002P1
  CURRENT APPLICATION NUMBER: US/09/149,476
  CURRENT FILING DATE: 1998-09-08
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  EARLIER FILING DATE: 1998-03-06
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  EARLIER FILING DATE: 1997-03-07
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EARLIER APPLICATION NUMBER: 60/043,669
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; EARLIER FILING DATE: 1997-08-22

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; EARLIER FILING DATE: 1997-08-22

; EARLIER APPLICATION NUMBER: 60/056,882

EARLIER FILING DATE: 1997-08-22

EARLIER APPLICATION NUMBER: 60/056,637

EARLIER FILING DATE: 1997-08-22

; EARLIER APPLICATION NUMBER: 60/056,903

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; EARLIER APPLICATION NUMBER: 60/056,888

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; EARLIER FILING DATE: 1997-08-22

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; EARLIER APPLICATION NUMBER: 60/056,911

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EARLIER FILING DATE: 1997-05-23

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; EARLIER APPLICATION NUMBER: 60/047,590

; EARLIER FILING DATE: 1997-05-23

; EARLIER APPLICATION NUMBER: 60/047,594

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   EARLIER APPLICATION NUMBER: 60/043,576
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   EARLIER APPLICATION NUMBER: 60/047,501
   EARLIER FILING DATE: 1997-05-23
   EARLIER APPLICATION NUMBER: 60/043,670
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   EARLIER APPLICATION NUMBER: 60/056,632
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   EARLIER APPLICATION NUMBER: 60/056,664
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  EARLIER APPLICATION NUMBER: 60/057,669
  EARLIER FILING DATE: 1997-09-05
  EARLIER APPLICATION NUMBER: 60/049.610
  EARLIER FILING DATE: 1997-06-13
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; Sequence 3, Application US/08700607
 Patent No. 5858708
  GENERAL INFORMATION:
    APPLICANT: Bandman, Olga
    APPLICANT: Au-Young, Janice
    APPLICANT: Goli, Surya K.
    APPLICANT: Hillman, Jennifer L.
    TITLE OF INVENTION: TWO NOVEL HUMAN NSP-LIKE PROTEINS
    NUMBER OF SEQUENCES: 9
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Incyte Pharmaceuticals, Inc.
      STREET: 3174 Porter Drive
      CITY: Palo Alto
      STATE: CA
      COUNTRY: U.S.
      ZIP: 94304
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette
      COMPUTER: IBM Compatible
      OPERATING SYSTEM: DOS
      SOFTWARE: FastSEQ Version 1.5
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/700,607
      FILING DATE: Filed Herewith
    ATTORNEY/AGENT INFORMATION:
      NAME: Billings, Lucy J.
      REGISTRATION NUMBER: 36,749
      REFERENCE/DOCKET NUMBER: PF-0114 US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 415-855-0555
      TELEFAX: 415-845-4166
  INFORMATION FOR SEQ ID NO: 3:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 241 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: peptide
    IMMEDIATE SOURCE:
      LIBRARY: THP1NOB01
      CLONE: 31870
US-08-700-607-3
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 Best Local Similarity 59.4%; Pred. No. 1.1e-25;
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              Db
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RESULT 8
US-08-714-741-32
; Sequence 32, Application US/08714741
; Patent No. 6500613
  GENERAL INFORMATION:
    APPLICANT: Briles, David E.
    APPLICANT: McDaniel, Larry S.
    APPLICANT: Swiatlo, Edwin
               Yother, Janet
    APPLICANT:
    APPLICANT: Crain, Marilyn J.
    APPLICANT: Hollingshead, Susan
    APPLICANT:
               Tart, Rebecca
    APPLICANT: Brooks-Walter, Alexis
    TITLE OF INVENTION: PNEUMOCOCCAL GENES, PORTIONS THEREOF,
    TITLE OF INVENTION: EXPRESSION PRODUCTS THEREFROM, AND USES OF SUCH GENES,
    TITLE OF INVENTION: PORTIONS AND PRODUCTS
    NUMBER OF SEQUENCES: 47
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Curtis, Morris & Safford, P.C.
      STREET: 530 Fifth Avenue
      CITY: New York
      STATE: New York
      COUNTRY: U.S.
      ZIP: 10036
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/714,741
      FILING DATE: 16-SEP-1996
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Frommer Esq., William S.
      REGISTRATION NUMBER: 25,506
      REFERENCE/DOCKET NUMBER: 454312-2460
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 840-3333
      TELEFAX: (212) 840-0712
  INFORMATION FOR SEQ ID NO: 32:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 8991 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
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; MOLECULE TYPE: amino acid US-08-714-741-32

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Qy 307	TKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVR	363
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Qy 553	KVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAP	596
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            Db
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; Patent No. 6420526
; GENERAL INFORMATION:
; APPLICANT: Rosen et al.
 TITLE OF INVENTION: 186 Human Secreted proteins
; FILE REFERENCE: PZ002P1
  CURRENT APPLICATION NUMBER: US/09/149,476
  CURRENT FILING DATE: 1998-09-08
  EARLIER APPLICATION NUMBER: PCT/US98/04493
  EARLIER FILING DATE: 1998-03-06
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; EARLIER FILING DATE: 1997-05-23

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; EARLIER FILING DATE: 1997-04-11
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- EARLIER APPLICATION NUMBER: 60/043,313
- EARLIER FILING DATE: 1997-04-11
- EARLIER APPLICATION NUMBER: 60/043,672
- EARLIER FILING DATE: 1997-04-11
- ; EARLIER APPLICATION NUMBER: 60/043,315
- EARLIER FILING DATE: 1997-04-11
- EARLIER APPLICATION NUMBER: 60/048,974
- ; EARLIER FILING DATE: 1997-06-06
- ; EARLIER APPLICATION NUMBER: 60/056,886
- ; EARLIER FILING DATE: 1997-08-22
- ; EARLIER APPLICATION NUMBER: 60/056,877
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- ; EARLIER FILING DATE: 1997-08-22

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EARLIER FILING DATE: 1997-08-22
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EARLIER FILING DATE: 1997-09-05

EARLIER FILING DATE: 1997-06-13

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; Sequence 8, Application US/08973462B
; Patent No. 6191270
; GENERAL INFORMATION:
  APPLICANT: DRUILHE, PIERRE
  APPLICANT: DAUBERSIES, PIERRE
  TITLE OF INVENTION: MALARIAL PRE-ERYTHROCYTIC STAGE POLYPEPTIDE MOLECULES
  FILE REFERENCE: 0660-0125-0 PCT
  CURRENT APPLICATION NUMBER: US/08/973,462B
  CURRENT FILING DATE: 1998-02-06
  EARLIER APPLICATION NUMBER: PCT/FR96/00894
  EARLIER FILING DATE: 1996-06-12
  EARLIER APPLICATION NUMBER: FR 95/07007
  EARLIER FILING DATE: 1995-06-13
  NUMBER OF SEQ ID NOS: 29
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   ORGANISM: Artificial Sequence
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; Sequence 5, Application US/08769309A
; Patent No. 5741890
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    APPLICANT: Scott, John D.,
    APPLICANT: Nauert, Brian J.,
    APPLICANT: Klauck, Theresa M.
    TITLE OF INVENTION: Protein Binding Domains of Gravin
    NUMBER OF SEQUENCES: 24
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: Marshall, O'Toole, Gerstein, Murray & Borun
      STREET: 6300 Sears Tower/233 South Wacker Drive
      CITY: Chicago
      STATE: Illinois
      COUNTRY: United States of America
      ZIP: 60606-6402
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/769,309A
      FILING DATE:
     CLASSIFICATION: 435
   ATTORNEY/AGENT INFORMATION:
    NAME: No. 5741890and, Greta E.
      REGISTRATION NUMBER: 35,302
      REFERENCE/DOCKET NUMBER:
                              27866/33451
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 312-474-6300
      TELEFAX: 312-474-0448
      TELEX: 25-3856
  INFORMATION FOR SEQ ID NO: 5:
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US-08-994-570-5
; Sequence 5, Application US/08994570
; Patent No. 6090929
 GENERAL INFORMATION:
   APPLICANT: Scott, John D.,
   APPLICANT: Nauert, Brian J.,
   APPLICANT: Klauck, Theresa M.
   TITLE OF INVENTION: Protein Binding Domains of Gravin
   NUMBER OF SEQUENCES: 24
   CORRESPONDENCE ADDRESS:
     ADDRESSEE: Marshall, O'Toole, Gerstein, Murray & Borun
     STREET: 6300 Sears Tower/233 South Wacker Drive
    CITY: Chicago
     STATE: Illinois
    COUNTRY: United States of America
    ZIP: 60606-6402
```

```
COMPUTER READABLE FORM:
     MEDIUM TYPE: Floppy disk
      COMPUTER: IBM PC compatible
     OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.30
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/994,570
     FILING DATE:
     CLASSIFICATION:
    ATTORNEY/AGENT INFORMATION:
     NAME: No. 6090929and, Greta E.
     REGISTRATION NUMBER: 35,302
     REFERENCE/DOCKET NUMBER: 27866/33451
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 312-474-6300
     TELEFAX: 312-474-0448
     TELEX: 25-3856
  INFORMATION FOR SEQ ID NO: 5:
    SEQUENCE CHARACTERISTICS:
     LENGTH: 1780 amino acids
     TYPE: amino acid
     TOPOLOGY: linear
    MOLECULE TYPE: protein
US-08-994-570-5
 Query Match
                    4.5%; Score 265.5; DB 3; Length 1780;
 Best Local Similarity 20.8%; Pred. No. 3e-08;
 Matches 264; Conservative 146; Mismatches 486; Indels 371; Gaps
         11 SSSTDSPPRPPPA----FKYQFVTE-----PEDE----EDEEEEEDEEED 47
Qу
           277 SKSAESPTSPVTSETGSTFK-KFFTQGWAGWRKKTSFRKPKEDEVEASEKKKEQEPEKVD 335
         48 DEDLEELEVLERKPAAGLSAAAVPPAAAA--PLLDFSSDSVPPAPRGPLPAAPPAAPERQ 105
Qу
            336 TEEDGKAEVASEKLTASEQAHPQEPÄESAHEPRLSAEYEKV-----ELPS-----EEO 383
        106 PSWERSPAAPAPSLPPAAAVLPSKLP-----EDDEPPARPPPPPPAGASP 150
           | : |: |: | : | |: |
Db
        384 VSGSQGPSEEKPA-PLATEVFDEKIEVHQEEVVAEVHVSTVEERTEEQKTEVEETAGSVP 442
        151 L-----AEP--AAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQP 201
QУ
                Dh
        443 AEELVGMDAEPQEAEPAKELVKLKETCVSGEDPTOGADLSPDEKVLSKPPEGVVSEVEML 502
Qу
        202 GNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLG-----NLSAVSSSEGTI 253
           503 SSQERMKVQGSPLKKLFTSTGLKKLS----GKKQKGKRGGGDEESGEHTQVPADSPDSQ 557
Db
        254 EETLNEASKELPERATN-PFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVI 312
Qу
           Db
        558 EEQKGESSASSPEEPEEITCLEKGLAEVQQDGEAEEGATSDGEKKREGVTPWASFKKMVT 617
        313 VRSK-----DKE---DLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMOMSVVAPV 362
Qу
                   :: []: []
        618 PKKRVRRPSESDKEDELDKVKSATLSSTEST------ASEMQEEMKGSVEEPK 664
Db
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	Qу	363	REEYADFKPFEQAWEVKDTYEGS	385
	Db	665	: : PEEPKRKVDTSVSWEALICVGSSKKRARRRSSSDEEGGPKAMGGDHQKADEAGKDKETGT	724
	Qу	386	RDVLAARANVESKVDRKCLEDSLEQKSLGKDS	417
	Db	725	: : : : : DGILAGSQEHDPGQGSSSPEQAGSPTEGEGVSTWESFKRLVTPRKKSKSKLEEKSEDS	782
	Qу	418	-EGRNEDASFPSTPEPVKDSS	453
	Db	783	IAGSGVEHSTPDT-EPGKEESWVSIKKFIPGRRKKRPDGKQEQAPVEDAGPTGANEDDSD	841
	Qу	454	TANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQD	501
	Db	842	VPAVVPLSEYDAVEREKMEAQQAQKGAEQPEQKAATEVSKELSESQVHMMAAAVADGTRA	901
	Qу	502		526
	Db	902	ATIIEERSPSWISASVTEPLEQVEAEAALLTEEVLEREVIAEEEPPTVTEPLPENR	957
-	Qy	527	LTPDLVQEA-CESELNEATGTKI-AYETKVDLVQTSEAIQESLYPTAQ	572
	Db	958	EARGDTVVSEAELTPEAVTAAETAGPLGSEEGTEASAAEETTEMVSAVSQLTDSPDTTEE	1017
	Qу	573	LCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVS	623
	Db	1018	ATPVQEVEGGVPDI EEQERRTQEVLQAVAEKVKEESQLPGTGGPEDVLQPVQ	1069
	Qy	624	YDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQE :	666
	Db	1070	RAEAERPEEQAEASGLKKETDVVLKVDAQEAKTEPFTQGKVVGQTTPESFEKAPQV	1125
	Qy	667	TEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPV	722
	Db	1126	TESPDSVETPT	1166
	Qy	723	DLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPY	774
	Db	1167	DSETDGSTPVADFDAPGTTQKDEIVEIHEENEVHLVPVRGTEAEAVPAQKERPP	1220
	Qy	775	LESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFS	834
	Db	1221	APSSFVFQEETKEQSKMEDTLEHTDKEVSVETVSILSKTEGTQEADQYA	1269
	Qу	835	DSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPC	885
	Db	1270	DEKTKDVPFFEGLEGSIDTGITVSREKVTEVALKGEGTEEAECKKDDALELQSHAKSPPS	1329
	Qу	886	LELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTE : : :: : : :	934
	Db	1330	PVEREMVVQVEREKTEAEPTHVNEEKLEHETAVTVSEEVSKQLLQTVNVPIIDGAKE	1386
	Qy	935	MGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVF: : : : : :	992
	Db	1387	VSSLEGSPPPCLGQEEAVCTKIQVQSSEASFTLTAAAEEEKVLGETANILETGETLEP	1444
	Qy	993	-GASLFL 998	

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RESULT 13
US-09-134-001C-4463
; Sequence 4463, Application US/09134001C
; Patent No. 6380370
; GENERAL INFORMATION:
  APPLICANT: Lynn Doucette-Stamm et al
  TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO
STAPHYLOCOCCUS
  TITLE OF INVENTION: EPIDERMIDIS FOR DIAGNOSTICS AND THERAPEUTICS
  FILE REFERENCE: GTC-007
  CURRENT APPLICATION NUMBER: US/09/134,001C
  CURRENT FILING DATE: 1998-08-13
  PRIOR APPLICATION NUMBER: US 60/064,964
  PRIOR FILING DATE: 1997-11-08
  PRIOR APPLICATION NUMBER: US 60/055,779
  PRIOR FILING DATE: 1997-08-14
  NUMBER OF SEQ ID NOS: 5674
 SEQ ID NO 4463
   LENGTH: 2137
   TYPE: PRT
   ORGANISM: Staphylococcus epidermidis
US-09-134-001C-4463
 Query Match
                      4.5%; Score 265; DB 4; Length 2137;
 Best Local Similarity 19.6%; Pred. No. 4.3e-08;
 Matches 172; Conservative 141; Mismatches 451; Indels 114; Gaps
                                                               24:
        159 STPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLE 218
Qу
            Db
        990 STSTSTSDSASTSTSE-----SESDSASTSLSESTSTSVSDSTSTSTSDSASMSASESE 1043
        219 TAASLPSLSPLSTVSFKEHGYLGNLSAV---SSSEGTIEETLNEASKELPERATNPFVNR 275
Qу
            1044 SNSKSTSLSESTSTSLS-----GSTSASTSDSASTSTSESESDSTSTSLSESTSTSLSGS 1098
Db
        276 DLAEFSELEYSEMGSSFKGSPKGESAILVE-----NTKEEVIVRSKDKEDLVC 323
Qy
             : | | : : |
       1099 TSASTSD---SASTSTSESDSTSESTSLSESLSTSVSDSTSASTSESASTSTSESESNSA 1155
Db
-Qу
        324 SAALHSPQESPVGKEDRVVSPEKTMDIFNEMOMSVVAPVREEYADFKPFEOAWEVKDTYE 383
            1156 STSLSGSLSTSISDSTSTSTSDSASTSTSESESDSTSTSLSE-----STSTSLSDSTS 1208
Db
Qу
        384 GSRDVLAARANVESKVDRKCLEDSLEQK---SLGKDSEGRNEDASFPSTPEPVKDSSRAY 440
               Db
       1209 TSTSESASTSTSES--DSTSESTSLSESTSTSVSDSTSASTSDSASTSTSVSDSESASTS 1266
Qу
        441 ITCASFTSATESTTANTFPLLEDHTSE--NKTDEKKIEERKAQIITEKTSPKTSNPFLVA 498
           |: : || ::||: :|
                               1267 ISESLSTSVSDSTSTSTSDSASTSTSESDSTSESTSLSESISTSVSDSTSASTSDSASTS 1326
Db
Qу
        499 VQDSEADYVTTD-----TLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYE 551
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1327 TSESESDSASTSLSGSTSTSLSDSTSTSTSDSASTSTSESDSERASTSLSGSTSTSLSDS 1386
Db
        552 TKV---DLVQTSEAIQESLYPTAOLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASV 608
Qу
          1387 TSTSTSDSASTSTSVSDS-----NSASTSLSGSLSTSVSDSTSTSTSDSASAST 1435
Db
        609 VQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALK-ALGTKEGIKEPESFNAAVQET 667
Qу
            1436 ---SESDSERA-----STSLSGSTSTSISDSTSTSTSDSASTSTSVSESNSTSTSISES 1486
Db
        668 EAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSD 727
Qу
                   1487 LSTSVS-----DSTSTSTSDSASTSTSVSDSDSASTSSSESV--STSDSESTSTSTS 1536
Db
        728 DSI---PEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQE----LGKPYLESFO 779
Qу
               Db
       1537 DSASTSTSVSESNSTSTSLSGSTSTSVSDSTSTSTSDSASASTSESDSDSASTSSSESVS 1596
QУ
       780 PNLH----STKDAASNDIPTLTKKEKISLOMEEFNTAIYSNDDLLSSKEDKIKESETFS 834
           1597 TSVSDSTSASTSESASTSTSVSDSNSASTSLSESTSTSLSDSTSMSTSDSASTSTSESDS 1656
Db
       835 DSSPIEIID-----EFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIOSGADS 882
QУ
           Dh
       1657 DSASTSLSDSTSTSVSESTSTSTSTSVSASNSTSTSLSDSRSTSLSDSTSTSTSESGSTS 1716
        883 LPCLELPCDLSFKNIYPKDEVHVSDEFSENR--SSVSKASISPSNVSALEPQTEMGSIVK 940
Qу
             1717 TS--ESDSDSASTSLSESTSTSISDSTSTSTSDSASTSMSVSDSNRASTSLSDSTSTSVS 1774
Db
Qу
       941 SKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVD 978
           Db
       1775 DSTSASTSESASTSTRESESTSASTSLS-ESTSTSVSD 1811
RESULT 14
US-08-978-277A-4
; Sequence 4, Application US/08978277A
; Patent No. 6582956
 GENERAL INFORMATION:
   APPLICANT: Gelman, Irwin H.
   TITLE OF INVENTION: TUMOR SUPPRESSOR GENE
   NUMBER OF SEQUENCES: 20
   CORRESPONDENCE ADDRESS:
     ADDRESSEE: Brumbaugh, Graves, Donohue & Raymond
     STREET: 30 Rockefeller Plaza
     CITY: New York
     STATE: NY
     COUNTRY: USA
     ZIP: 10112-0228
   COMPUTER READABLE FORM:
     MEDIUM TYPE: Diskette
     COMPUTER: IBM Compatible
     OPERATING SYSTEM: DOS
    SOFTWARE: FastSEQ Version 1.5
   CURRENT APPLICATION DATA:
    APPLICATION NUMBER: US/08/978,277A
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FILING DATE:
     CLASSIFICATION: 514
    PRIOR APPLICATION DATA:
     APPLICATION NUMBER: 08/665,401
     FILING DATE: 18-JUN-1996
    ATTORNEY/AGENT INFORMATION:
     NAME: Clark, Richard S
     REGISTRATION NUMBER: 26,154
     REFERENCE/DOCKET NUMBER: A30558 - 165/34008
    TELECOMMUNICATION INFORMATION:
     TELEPHONE: 212-408-2558
     TELEFAX: 212-765-2519
     TELEX:
  INFORMATION FOR SEO ID NO: 4:
    SEQUENCE CHARACTERISTICS:
     LENGTH: 1596 amino acids
     TYPE: amino acid
     STRANDEDNESS: single
     TOPOLOGY: linear
    MOLECULE TYPE: peptide
    HYPOTHETICAL: NO
    ANTI-SENSE: NO
    FRAGMENT TYPE: internal
    ORIGINAL SOURCE:
US-08-978-277A-4
 Query Match
                    4.4%; Score 254.5; DB 4; Length 1596;
 Best Local Similarity 19.9%; Pred. No. 1.3e-07;
 Matches 258; Conservative 162; Mismatches 468; Indels 409; Gaps
         7 SSLVSSSTDSP-----PRPPPAFKYQFVTEPEDEEDEEEEDDEDLEELE 55
Qу
           Db
        273 SSPVNSETTSSFKKFFTHGWAGWRKKTSFKKSKEDDLETAEKRKEOEAEKVDEEEKEKTE 332
        56 VL--ERKPAAGLSAAAVP---PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWER 110
Qу
             333 PASEEQEPAEDTDQARLSADYEKVELPLEDOVGDLEASSEEKCAPLATEVFDEKMEAHOE 392
QУ
        111 SPA----- 135
                                      Db
        393 VVAEVHVSTVEKTEEEQGGGGEAEGGVVVEGTGESLPPEKLAEPQEVPQEAEPAEELMKS 452
        136 ------ASPLAEPAAPP--- 158
Qу
                        1 : ' | | |
                                               Db
        453 REMCVSGGDHTQLTDLSPEEKTLPKHPEGIVSEVEMLSSQERIKVQGSPLKKLFSSSGLK 512
        159 -- STPAAPKRRGSGSVDETLFALPAASEPV---IPSSAEKIMDLMEQPG-NTVSSGQEDF 212
Qу
            513 KLSGKKQKGKRGGGGDEE----PGEYQHIHTESPESAD-----EQKGESSASSPEEPE 561
Dh
Qу
        213 PSVLLET-----
                                    ----AASLPSLSPLSTV----SFKEHG 238
           : []
                                       |: ::| | | | |
        562 ETTCLEKGPLEAPQDGEAEEGTTSDGEKKREGITPWASFKKMVTPKKRVRRPSESDKEEE 621
Db
Qу
        239 YLGNLSA-VSSSEGTIEE-----TLNEASK-ELPERATNPFVNRD----LAEFSELEYS 286
               622 LEKVKSATLSSTDSTVSEMQDEVKTVGEEQKPEEPKRRVDTSVSWEALICVGSSKKRARK 681
Db
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Qу	287	EMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKE	338
Db	682	ASSSDDEGGPRTLGGDSHRAEEASKDKEAGTDAVPASTQEQDQAQGSSSPEPAGSPSEGE	741
Qу	339	DRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAW-EVKD	380
Db	742	GVSTWESFKRLVTPRKKSKSKLEEKAEDSSVEQLSTEIEPSREESWVSIKK	792
Qy	381	TYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDS	417
Db	793	FIPGRRKKRADGKQEQATVEDSGPVEINEDDPNVPAVVPLSEYNAVEREKMEA	845
Qy	418	EGRNEDASFPSTPEPVKDSSRAYITCASFTSAT:	450
Db	846	QGNTELPQLLGAVYVSEELSKTLVHTVSVAVIDGTRAVTSVEERSPSWIS-ASVTEPL	902
Qy	451	ESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTD	510
Db	903	EHTAGEAMPPVEEVTEKDIIAEETPVLTQ-TLPEGKDAHDDMVTSE	947
Qу	511	TLSKVTEAAVSNMPEGLTPDLVQEACES-ELNEATGTKIAYETKVDLVQTSEAIQESLYP	569
Db	948		994
Qy	570	TAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPP	620
Db	995	TEEATPVQEVESGVLDTEEEERQTQAILQAVADKVKEESQVP-ATQTVQRTGSKALEKVE	1053
Qy	621	PVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISI	674
Db	1054	EVEEDSEVLASEKEKDVMPKGPVQEAGAEHL-AQGSETGQATPESLEVPEVTADVDHVA-	1111
Qу	675	ACDLIKETKLSTEP-SPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDS	729
Db	1112	TCQVIKLQQLMEQAVAPESSETLTDSETNGSTPLADSDTADGTQQDETIDSQDSKATA	1169
Qу	730	-IPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDA : : :	788
Db	1170	AVRQSQVTEEEAATAQKEEPSTLPNNVPAQEEHGEEPGRDVLEPTQQELTA	1220
Qy	789	ASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLS : : : : :	821
Db	1221	AAVPVLAKTEVGQEGEVDWLDGEKVKEEQEVFVHSGPNSQKAADVTYDSEVMGVAGCQ	1278
Qy	822	SKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDL : : : : : : : : :	865
Db	1279	EKESTEVQSLSLEEGEMETDVEK-EKRETKPEQVSEEGEQETAAPEHEGTYGKPVLTLDM	1337
QУ	866	EVSDKSEIANIQSGADSLPCLE	887
Db	1338	PSSERGKALGSLGGSPSLPDQDKAGCIEVQVQSLDTTVTQTAEAVEKVIETVVISETGES	1397
Qу	888	LPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMG	936
Db	1398	PECVGAHLLPAEKSSATGGHWTLOHAEDTVPLGPESOAESIPIIVTPAPESTLHPDLO-G	1456

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937 SIVKSKSLTKEAEKKL---PSDTEKEDRSLSAVLSAE 970
Qу
             Db
        1457 EISASQRERSEEEDKPDAGPDADGKESTAIEKVLKAE 1493
RESULT 15
US-08-993-118-7
; Sequence 7, Application US/08993118
; Patent No. 5997872
  GENERAL INFORMATION:
    APPLICANT: LUCAS, Sophie;
    APPLICANT: DE SMET, Charles;
    APPLICANT: BOON-FALLEUR, Thierry
    TITLE OF INVENTION: ISOLATED NUCLEIC ACID MOLECULE CODING FOR TUMOR
    TITLE OF INVENTION: REJECTION ANTIGEN PRECURSOR MAGE-C1 AND USES
    TITLE OF INVENTION: THEREOF
    NUMBER OF SEQUENCES: 14
    CORRESPONDENCE ADDRESS:
     ADDRESSEE: Felfe & Lynch
      STREET: 805 Third Avenue
      CITY: New York City
     STATE: New York
     COUNTRY: USA
     ZIP: 10022
    COMPUTER READABLE FORM:
      MEDIUM TYPE: Diskette, 3.5 inch, 360 kb storage
      COMPUTER: IBM PS/2
      OPERATING SYSTEM: PC-DOS
      SOFTWARE: Wordperfect
    CURRENT APPLICATION DATA:
      APPLICATION NUMBER: US/08/993,118
      FILING DATE:
      CLASSIFICATION:
    PRIOR APPLICATION DATA:
      APPLICATION NUMBER: US/08/845,528
      FILING DATE: April 25, 1997
    ATTORNEY/AGENT INFORMATION:
      NAME: Mary Anne Schofield
      REGISTRATION NUMBER: 36,669
      REFERENCE/DOCKET NUMBER: LUD 5455
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: (212) 688-9200
      TELEFAX: (212) 838-3884
  INFORMATION FOR SEQ ID NO: 7:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 1142
      TYPE: amino acids
      STRANDEDNESS: single-stranded
      TOPOLOGY: linear
US-08-993-118-7
 Query Match
                        4.0%; Score 233; DB 2; Length 1142;
 Best Local Similarity 19.6%; Pred. No. 1.9e-06;
 Matches 221; Conservative 156; Mismatches 439; Indels 310; Gaps
           8 SLVSSSTDSPPRPPPAFKYQFVTEPEDEEDE-EEEEDEEEDDEDLEELEVLERKPAAGLS 66
Qу
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	Db	13	SLLQSSSESPQSCPEGEDSQSPLQIPQSSPESDDTLYPLQSPQSRSEG	60	
	Qy	67	AAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLPPAAA : : : : : : :	124	
	Db	61	: : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : :	105	
	ДУ	125	VLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGS :	171	
	Db	106	KDSLSPLEISQSPPEGEDVQSPLQNPASSFFSSALLSIFQSSPESIQSPFEG-	157	
	Qy	172	VDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLE	218	
	Db	158	FPQSVLQIPVSAASSTLVSIFQSSPESTQSPFEGFPQSPLQIPVSRSFSSTLLS	212	
	Qу	219	TAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLA	278	
	Db	213	IFQSSPERSQRTSEGFAQSPLQIPVSSSSSSTLLSLFQSSPERTQ	257	
	QУ	279	EFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESP	334	
	Db	258		306	
	QУ	335	VGKEDRVVSPEKTMDIFNEMQMSVV-APVREEYADFKPFEQAWE-VKDTYE	383	-
	Db	307	VSSSSSTLLSLFQSSPERTQSTFEGFPQSLLQIPMTSSFSSTLLSIFQSSPESAQSTFE	366	
. *	Qу	384	GSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITC	443	
	Db	367	GFPQSPLQIPGSPSFSSTL	385	
	Qу	444	ASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLV :: : : :: : : :: ::	497	
	Db	386	LSLFQSSPERTHSTFEGFPQSPLQIPMTSSFSSTLLSILQSSPESAQSAFE	436	
	QУ	498	AVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLV	557	
	Db	437	GFPQSPLQIPVSSSFSYTLLSLFQSSPE-RTQSTFEGFPQSPLQIPVSSSSSSSTLLSLF	495	+
	QУ	558	QTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPS	603	
	Db		QSSPECTQSTFEGFPQSPLQIPQSPPEGENTHSPLQIVPSLPEWEDSLSPHYFPQSPP		
	QУ		AGASVVQPSVSPLEAPPPVSYDSIKLE-PENPPPYEEAMNVALKA		
	Db		QGEDSLSPHYFP-QSPPQGEDSLSPHYFPQSPQGEDSLSPHYFPQSPPQGEDSMSPLYFP		
	Qу		LGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPE : :: :: ::		
	Db		QSPLQGEEFQSSLQSPVSI-CSSSTPSSLPQSFPESSQSPPEGPV		
	Qy		HAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : :		
	Db		QSPLHSPQSPPEGMHSQSPLQSPESAPEGEDSLSPLQIPQSPLEGEDSLSSLH		
	ДУ		ETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKIS : : : : : :		
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Search completed: December 19, 2003, 15:37:55 Job time : 25 secs

GenCore version 5.1.6 Copyright (c) 1993 - 2003 Compugen Ltd.

OM protein - protein search, using sw model

Run on:

December 19, 2003, 15:36:49; Search time 40 Seconds

(without alignments)

5430.007 Million cell updates/sec

Title:

US-09-830-972-2

Perfect score:

5848

Sequence:

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Total number of hits satisfying chosen parameters:

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Maximum DB seq length: 2000000000

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Maximum Match 100%

Listing first 45 summaries

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Published Applications AA:*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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Result

Query

No. Score Match Length DB

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ALIGNMENTS

RESULT 1

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- ; Patent No. US20020072493A1
- ; GENERAL INFORMATION:
- ; APPLICANT: EISENBACH-SCHWARTZ, Michal

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APPLICANT: COHEN, Irun R.
  APPLICANT:
            BESERMAN, Pierre
  APPLICANT: MOSONEGO, Alon
  APPLICANT:
            MOALEM, Gila
  TITLE OF INVENTION: ACTIVATED T-CELLS, NERVOUS SYSTEM-SPECIFIC ANTIGENS AND
  FILE REFERENCE: EIS-SCHWARTZ=2A
  CURRENT APPLICATION NUMBER: US/09/893,348
  CURRENT FILING DATE: 2001-06-28
  PRIOR APPLICATION NUMBER: US 09/314,161
  PRIOR FILING DATE: 1999-05-19
  PRIOR APPLICATION NUMBER: US 09/218,277
  PRIOR FILING DATE: 1998-12-22
  PRIOR APPLICATION NUMBER: PCT/US98/14715
  PRIOR FILING DATE: 1998-07-21
  PRIOR APPLICATION NUMBER: IL 124500
  PRIOR FILING DATE: 1998-05-19
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US-09-758-140-6

[;] Sequence 6, Application US/09758140 ; Patent No. US20020012965A1

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  APPLICANT: Strittmatter, Stephen M.
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Axonal Growth
  FILE REFERENCE: 44574-5073-US
  CURRENT APPLICATION NUMBER: US/09/758.140
  CURRENT FILING DATE: 2001-01-12
  PRIOR APPLICATION NUMBER: US 60/175,707
  PRIOR FILING DATE: 2000-01-12
  PRIOR APPLICATION NUMBER: US 60/207,366
  PRIOR FILING DATE: 2000-05-26
  PRIOR APPLICATION NUMBER: US 60/236,378
  PRIOR FILING DATE: 2000-09-29
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   ORGANISM: Homo sapiens
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US-09-972-599A-6

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- ; GENERAL INFORMATION:
- ; APPLICANT: STRITTMATTER, STEPHEN M.
- ; TITLE OF INVENTION: NOGO-RECEPTOR-MEDIATED BLOCKADE OF AXONAL GROWTH
- ; FILE REFERENCE: C077 CIP US

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  CURRENT FILING DATE: 2001-10-06
  PRIOR APPLICATION NUMBER: PCT/US01/01041
  PRIOR FILING DATE: 2001-01-12
  PRIOR APPLICATION NUMBER: 09/758,140
  PRIOR FILING DATE: 2001-01-12
  PRIOR APPLICATION NUMBER: 60/236,378
  PRIOR FILING DATE: 2000-09-29
  PRIOR APPLICATION NUMBER: 60/207,366
  PRIOR FILING DATE: 2000-05-26
  PRIOR APPLICATION NUMBER: 60/175,707
  PRIOR FILING DATE: 2000-01-12
  NUMBER OF SEQ ID NOS: 57
  SOFTWARE: PatentIn Ver. 2.1
 SEQ ID NO 6
   LENGTH: 1192
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-972-599A-6
 Query Match
                     75.3%; Score 4403.5; DB 9; Length 1192;
 Best Local Similarity 75.9%; Pred. No. 4.7e-214;
 Matches 909; Conservative 104; Mismatches 145; Indels
                                                    39; Gaps
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Qу	869	DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	926
Db	896	HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	955
Qу	927	SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	986
Db	956	:	1015
Qу	987	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1046
Db	1016	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1075
Qу	1047	YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1106
Db	1076	YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1135
Qу	1107	NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 116	3
Db	1136	NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 119	2

US-10-060-036-71

- ; Sequence 71, Application US/10060036
- ; Publication No. US20030073144A1
- ; GENERAL INFORMATION:
- ; APPLICANT: Benson, Darin R.
- ; APPLICANT: Kalos, Michael D.
- ; APPLICANT: Lodes, Michael J.
- ; APPLICANT: Persing, David H.

```
APPLICANT: Hepler, William T.
  APPLICANT: Jiang, Yuqiu
  TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE THERAPY
  TITLE OF INVENTION: AND DIAGNOSIS OF PANCREATIC CANCER
  FILE REFERENCE: 210121.566
  CURRENT APPLICATION NUMBER: US/10/060.036
  CURRENT FILING DATE:
                   2002-01-30
  NUMBER OF SEQ ID NOS: 4560
  SOFTWARE: FastSEQ for Windows Version 4.0
 SEQ ID NO 71
   LENGTH: 1192
   TYPE: PRT
   ORGANISM: Homo sapiens
US-10-060-036-71
 Query Match
                    75.3%; Score 4403.5; DB 15; Length 1192;
 Best Local Similarity
                    75.9%; Pred. No. 4.7e-214;
 Matches 909; Conservative 104; Mismatches 145; Indels
                                                  39; Gaps
                                                            20:
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
Qу
           Db
         1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDEDLEELEVLERK 58
         61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
Qу
           59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP 118
Db
        116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----
Qу
                                               -PAAPPSTPAAPKR 166
                Db
        119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPOAEPVWTPPAPAPAAPPSTPAAPKR 178
        167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Qу
           179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEQPGNTISAGQEDFPSVLLETAASLPS 237
Db
Qу
        226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
           238 LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY 297
Db
        286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVG----KED 339
Qу
           Db
        298 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNQQELPTALTKLVKED 357
        340 RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAA----RANV 395
Qу
                 358 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 416
Db
        396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
Qу
           Db
       417 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRSGAYITCAPFNPAATESIA 476
       455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS 513
Qу
            Db
       477 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 536
       514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
Qу
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Db	537	KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL	596
Qу	574	CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN	633
Db	597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN	655
Qy	634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF	692
Db	656	PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF	715
Qy		SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS	752
Db	716	SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS	775
Qу	753	-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF	809
Db	776	FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLQMEEL	835
Qу		NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS : :	
Db	836	STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS	895
Qу		DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	
Db		HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	
Qу		SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	
Db		SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK	
Qу		KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	
Db		KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1075
QУ			1106
Db	1076	YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1135
Qу		NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 116	
Db	1136	NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 119	12

US-09-789-386-2

- ; Sequence 2, Application US/09789386
- ; Patent No. US20020010324A1
- ; GENERAL INFORMATION:
- ; APPLICANT: MICHALOVICH, DAVID
- APPLICANT: PRINJHA, RABINDER KUMAR
- ; TITLE OF INVENTION: NOVEL COMPOUNDS
- FILE REFERENCE: GP-30165-C1
- ; CURRENT APPLICATION NUMBER: US/09/789,386
- ; CURRENT FILING DATE: 2001-02-21
- ; PRIOR APPLICATION NUMBER: U.K. 9916898.1
- ; PRIOR FILING DATE: 1999-07-19
- ; PRIOR APPLICATION NUMBER: U.K. 9816024.5

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PRIOR FILING DATE: 1998-07-22
  PRIOR APPLICATION NUMBER: US 09/359,208
  PRIOR FILING DATE: 1999-07-22
  NUMBER OF SEQ ID NOS: 6
  SOFTWARE: FastSEQ for Windows Version 3.0
 SEO ID NO 2
   LENGTH: 1192
   TYPE: PRT
   ORGANISM: HOMO SAPIENS
US-09-789-386-2
 Query Match
                    75.2%; Score 4398.5; DB 9; Length 1192;
 Best Local Similarity
                    75.9%; Pred. No. 8.4e-214;
 Matches 908; Conservative 104; Mismatches 146;
                                           Indels
                                                  39; Gaps
                                                           20;
Qу
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
           Db
         1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEEDEDEDLEELEVLERK 58
        61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
Qу
           59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP 118
Db
        116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
Qу
               Db
        119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAAPPSTPAAPKR 178
        167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Qу
           Db
        179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEQPGNTISAGQEDFPSVLLETAASLPS 237
        226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
Qу
           Db
        238 LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY 297
Qу
       286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVG----KED 339
           Db
       298 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNOOELPTALTKLVKED 357
       340 RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAA----RANV 395
Qу
           358 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 416
Db
Qу
       396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
           |||||:|| |||||
                             417 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRPGAYITCAPFNPAATESIA 476
Db
Qу
       455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS 513
           Db
       477 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 536
Qу
       514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
              Db
       537 KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL 596
       574 CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN 633
Qу
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Db	597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN	655
Qу	634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF	692
Db	656	PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF	715
Qy	693	SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS	752
Db	716	SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS	77Ś
Qу	753	-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF :: ::: :	809
Db	776	FESMI EYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKI PLQMEEL	835
Qy	810	NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS: :	868
Db	836	STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS	895
Qу	869	DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	926
Db	896	HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	955
Qу	927	SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	986
Db		SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK	
· Qy	987	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1046
Db	1016	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1075
Qу	1047	YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1106
Db	1076	YLESEVAÍSEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1135
Qу	1107	NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 116	53
Db	1136	NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 119	92

- ; Sequence 23, Application US/09893348
- ; Patent No. US20020072493A1
- ; GENERAL INFORMATION:
- ; APPLICANT: EISENBACH-SCHWARTZ, Michal
- ; APPLICANT: COHEN, Irun R.
- ; APPLICANT: BESERMAN, Pierre
- ; APPLICANT: MOSONEGO, Alon
- ; APPLICANT: MOALEM, Gila
- ; TITLE OF INVENTION: ACTIVATED T-CELLS, NERVOUS SYSTEM-SPECIFIC ANTIGENS AND THEIR USES
- ; FILE REFERENCE: EIS-SCHWARTZ=2A
- ; CURRENT APPLICATION NUMBER: US/09/893,348
- ; CURRENT FILING DATE: 2001-06-28
- ; PRIOR APPLICATION NUMBER: US 09/314,161
- ; PRIOR FILING DATE: 1999-05-19
- ; PRIOR APPLICATION NUMBER: US 09/218,277

```
PRIOR FILING DATE: 1998-12-22
  PRIOR APPLICATION NUMBER: PCT/US98/14715
  PRIOR FILING DATE: 1998-07-21
  PRIOR APPLICATION NUMBER: IL 124500
  PRIOR FILING DATE: 1998-05-19
  NUMBER OF SEQ ID NOS: 29
  SOFTWARE: PatentIn version 3.1
 SEQ ID NO 23
   LENGTH: 1192
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-893-348-23
 Query Match
                    75.2%; Score 4398.5; DB 9; Length 1192;
 Best Local Similarity
                    75.9%; Pred. No. 8.4e-214;
 Matches 908; Conservative 104; Mismatches 146; Indels
                                                  39; Gaps
                                                            20;
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
Qу
           1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDLEELEVLERK 58
Db
Qу
         61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
           59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPEROPSWDPSPVSSTVP 118
Db
        116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
Qу
                Db
        119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAPPSTPAAPKR 178
        167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
QУ
           179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEOPGNTISAGQEDFPSVLLETAASLPS 237
Db
        226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
Qу
           ::||| |||||
        238 LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY 297
Db
        286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVG----KED 339
Qу
                 298 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNQQELPTALTKLVKED 357
Dh
Qу
        340 RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEOAWEVKDTYEGSRDVLAA----RANV 395
            358 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 416
Db
        396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
QУ
           417 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRPGAYITCAPFNPAATESIA 476
Db
        455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS 513
Qу
            Dh
        477 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 536
        514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
Qy
               Dh
        537 KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAOL 596
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Qу	574	CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN	633
Db	597	CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN	655
Qy	634	PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF	692
Db	656	PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF	715
Qy	693	SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS	752
Db	716	SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS	775
Qу	753	-ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF :: ::: : : : : ::: :	809
Db	776	FESMI EYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKI PLQMEEL	835
Qу	810	NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS: :	868
Db	836	STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS	895
Qу	869	DKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNV	926
Db	896	HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV	955
Qy	927	SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK	986
Db	956	SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK	1015
Qy	987	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1046
Db	1016	KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA	1075
Qу	1047	YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF	1106
Db	1076	W. DOWN TOWN	1135
Qy -	1107	NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 116	53
Db	1136	NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 119	2

US-10-205-194-164

- ; Sequence 164, Application US/10205194
- ; Publication No. US20030134301A1
- ; GENERAL INFORMATION:
- ; APPLICANT: Warner-Lambert Company
- APPLICANT: Lee, Kevin
- ; APPLICANT: Dixon, Alistair
- ; APPLICANT: Brooksbank, Robert
- ; APPLICANT: Pinnock, Robert
- TITLE OF INVENTION: Identification and Use of Molecules Implicated in Pain
- ; FILE REFERENCE: WL-A-018201
- ; CURRENT APPLICATION NUMBER: US/10/205,194
- ; CURRENT FILING DATE: 5200-07-24
- ; PRIOR APPLICATION NUMBER: GB 0118354.0
- ; PRIOR FILING DATE: 2001-07-27

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; NUMBER OF SEQ ID NOS: 177
  SOFTWARE: PatentIn Ver. 2.1
; SEO ID NO 164
   LENGTH: 379
   TYPE: PRT
   ORGANISM: Rattus norvegicus
   OTHER INFORMATION: Foocen-m2 reticulon
US-10-205-194-164
 Query Match
                  25.9%; Score 1513; DB 12; Length 379;
 Best Local Similarity 32.6%; Pred. No. 8.1e-69;
 Matches 379; Conservative 0; Mismatches 0; Indels 784; Gaps
Qу
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
          Db
         1 MEDIDOSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
        61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120
Qу
          61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120
Db
Qу
       121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
          121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
Db
       181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240
Qу
          1111111111
       181 AASEPVIPSSA----- 191
Db
       241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300
Qу
       192 ----- 191
Db
Qу
       301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVA 360
Db
       Qу
       361 PVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEOKSLGKDSEGR 420
Db
       192 ----- 191
       421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480
Qу
Db
Qу
       481 QIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESEL 540
       192 ----- 191
Db
Qy
       541 NEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSL 600
Db
       192 ----- 191
Qу
       601 LPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESF 660
       192 ----- 191
Dh
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(ΣУ	661	NAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESE	720
]	Ob	192	·	191
(Qу	721	PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQP	780
-]	Ob	192		191
(ДУ	781	NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	840
]	Ob	192		191
(QY	841	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK	900
]	Db .	192		191
(ДУ	901	DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED	960
1	Ob	192		191
. (ДÄ	961	RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS	1020
I	Ob	192	VVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS	236
(Σλ "	1021	VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	1080
Ι	Ob	237	VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	296
(Ο̈́Α	1081	FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLA	1140
Ι	Ob	297	FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLA	356
(JA	1141	NKSVKDAMAKIQAKIPGLKRKAD 1163	
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- ; Sequence 20, Application US/09893348
- ; Patent No. US20020072493A1
- ; GENERAL INFORMATION:
- ; APPLICANT: EISENBACH-SCHWARTZ, Michal
- ; APPLICANT: COHEN, Irun R.
- ; APPLICANT: BESERMAN, Pierre
- ; APPLICANT: MOSONEGO, Alon
- ; APPLICANT: MOALEM, Gila
- ; TITLE OF INVENTION: ACTIVATED T-CELLS, NERVOUS SYSTEM-SPECIFIC ANTIGENS AND THEIR USES
- ; FILE REFERENCE: EIS-SCHWARTZ=2A
- ; CURRENT APPLICATION NUMBER: US/09/893,348
- CURRENT FILING DATE: 2001-06-28
- ; PRIOR APPLICATION NUMBER: US 09/314,161
- ; PRIOR FILING DATE: 1999-05-19
- ; PRIOR APPLICATION NUMBER: US 09/218,277
- ; PRIOR FILING DATE: 1998-12-22

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PRIOR APPLICATION NUMBER: PCT/US98/14715
  PRIOR FILING DATE: 1998-07-21
  PRIOR APPLICATION NUMBER: IL 124500
  PRIOR FILING DATE: 1998-05-19
  NUMBER OF SEQ ID NOS: 29
  SOFTWARE: PatentIn version 3.1
: SEO ID NO 20
   LENGTH: 360
   TYPE: PRT
   ORGANISM: Rattus norvegicus
US-09-893-348-20
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                    24.1%; Score 1411.5; DB 9; Length 360;
 Best Local Similarity 31.0%; Pred. No. 1e-63;
 Matches 360; Conservative 0; Mismatches 0; Indels 803; Gaps
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         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEDDEDLEELEVLERK 60
        61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120
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           Db
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        121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
Qу
           Db
        121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSV----- 172
Qу
        181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240
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Qу
       241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300
       173 ----- 172
Db
       301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVA 360
Qу
       173 ----- 172
Db
Qу
       361 PVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGR 420
Db
QУ
       421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480
Db
       173 ----- 172
Qу
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       173 ----- 172
Db
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Db
       173 ----- 172
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Db	173		172
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Db	173		172
Qу	781	NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	840
Db	173		172
Qу	841	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK	900
Db	173		172
Qу	901	DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED	960
Db	173		172
Qу	961	RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS	1020
Db	173		217
Qу	1021	VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	1080
Db	218	VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL	277
Qу	1081	FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLA	1140
Db	278	FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLA	337
Qу	1141	NKSVKDAMAKIQAKIPGLKRKAD 1163	
Db	338		

US-09-789-386-6

- ; Sequence 6, Application US/09789386
- ; Patent No. US20020010324A1
- ; GENERAL INFORMATION:
- APPLICANT: MICHALOVICH, DAVID
- ; APPLICANT: PRINJHA, RABINDER KUMAR
- ; TITLE OF INVENTION: NOVEL COMPOUNDS
- ; FILE REFERENCE: GP-30165-C1
- CURRENT APPLICATION NUMBER: US/09/789,386
- ; CURRENT FILING DATE: 2001-02-21
- PRIOR APPLICATION NUMBER: U.K. 9916898.1
- ; PRIOR FILING DATE: 1999-07-19
- ; PRIOR APPLICATION NUMBER: U.K. 9816024.5
- PRIOR FILING DATE: 1998-07-22
- ; PRIOR APPLICATION NUMBER: US 09/359,208
- ; PRIOR FILING DATE: 1999-07-22

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NUMBER OF SEQ ID NOS: 6
  SOFTWARE: FastSEQ for Windows Version 3.0
 SEQ ID NO 6
   LENGTH: 373
   TYPE: PRT
   ORGANISM: HOMO SAPIENS
US-09-789-386-6
  Query Match
                      20.4%; Score 1191; DB 9; Length 373;
  Best Local Similarity 27.8%; Pred. No. 1.4e-52;
 Matches 327; Conservative 12; Mismatches 20; Indels 818; Gaps
                                                                 7;
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            Db
          1 MEDLDQSPLVSSS-DSPPRPQPAFKYQFVREPEDEE-EEEEEEEEDEDEDLEELEVLERK 58
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            59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP 118
Db
        116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE-----PAAPPSTPAAPKR 166
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                Db
        119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
        167 RGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSL 226
QУ
            Db
Qу
        227 SPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYS 286
Db
Qу
        287 EMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEK 346
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        183 ----- 182
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        467 ENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEG 526
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        527 LTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSP 586
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        587 VLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALK 646
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Qу
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        767 PQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDK 826
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Db
        827 IKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCL 886
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        887 ELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPOTEMGSIVKSKSLTK 946
Qу
Db
        183 ------GSV------ 185
        947 EAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFS 1006
Qу
                                     Db
                                -----VVDLLYWRDIKKTGVVFGASLFLLLSLTVFS 216
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        1007 IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVOKYSNSA 1066
            Db
        217 IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVOKYSNSA 276
Qу
       1067 LGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIY 1126
            Db
        277 LGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSVPVIY 336
Qу
       1127 ERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
            337 ERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 373
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RESULT 10
US-09-765-205-6
; Sequence 6, Application US/09765205
; Patent No. US20020034800A1
; GENERAL INFORMATION:
  APPLICANT: Cao, Li
  TITLE OF INVENTION: BONE MARROW SECRETED PROTEINS AND POLYNUCLEOTIDES
  FILE REFERENCE: 1458.004/200130.449
  CURRENT APPLICATION NUMBER: US/09/765,205
  CURRENT FILING DATE: 2001-01-17
  PRIOR APPLICATION NUMBER: US/09/212,440
  PRIOR FILING DATE: 1998-12-16
  NUMBER OF SEQ ID NOS: 46
  SOFTWARE: FastSEQ for Windows Version 3.0
; SEO ID NO 6
   LENGTH: 373
   TYPE: PRT
   ORGANISM: human
US-09-765-205-6
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20.4%; Score 1191; DB 9; Length 373;

Query Match

		Similarity 27.8%; Pred. No. 1.4e-52; 7; Conservative 12; Mismatches 20; Indels 818; Gaps	7
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Db	59	PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP	118
Qу	116	APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKR	166
Db	119	APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR	178
Qу	167	RGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSL	226
Db	179	RGSS	182
Qу	227	SPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYS	286
Db	183		182
Qу	287	EMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEK	346
Db	183		182
Qу		TMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLED	
Db	183		182
Qу	407	SLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTS	466
Db	183		182
Qу	467	ENKTDEKKI EERKAQI I TEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEG	526
Db	183	·	182
Qу	527	LTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSP	586
Db	183		182
Qу	587	VLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALK	646
Db	183		182
Qу	647	ALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVP	706
Db	183		182
Qу	707	EHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSAS	766
Db	183		182
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	Db	183		182
	Qy	827	IKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCL	886
	Db	183		182
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	Db	183	: GSV	185
	Qу		EAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFS	
	Db	186		216
	Qу	1007	IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSA	1066
	Db	217	IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSA	276
	Qу	1067	LGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIY	1126
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	US-09-893			
; Sequence 24, Application US/09893348				
; Patent No. US20020072493A1 ; GENERAL INFORMATION:				
; APPLICANT: EISENBACH-SCHWARTZ, Michal				
	; APPLICANT: COHEN, Irun R.			
; APPLICANT: BESERMAN, Pierre ; APPLICANT: MOSONEGO, Alon				
; APPLICANT: MOSONEGO, ATOM ; APPLICANT: MOALEM, Gila				
; TITLE OF INVENTION: ACTIVATED T-CELLS, NERVOUS SYSTEM-SPECIFIC ANTIGENS AND				
THEIR USES				
	; FILE REFERENCE: EIS-SCHWARTZ=2A ; CURRENT APPLICATION NUMBER: US/09/893,348			
	; CURRENT FILING DATE: 2001-06-28			
	; PRIOR APPLICATION NUMBER: US 09/314,161			
			G DATE: 1999-05-19	

PRIOR APPLICATION NUMBER: US 09/218,277

PRIOR APPLICATION NUMBER: PCT/US98/14715

PRIOR APPLICATION NUMBER: IL 124500

PRIOR FILING DATE: 1998-12-22

PRIOR FILING DATE: 1998-07-21

PRIOR FILING DATE: 1998-05-19 NUMBER OF SEQ ID NOS: 29

SOFTWARE: PatentIn version 3.1

ORGANISM: Homo sapiens

SEQ ID NO 24 LENGTH: 373 TYPE: PRT

	Query Ma Best Loc		20.4%; Score 1191; DB 9; Length 373; Similarity 27.8%; Pred. No. 1.4e-52;	
			7; Conservative 12; Mismatches 20; Indels 818; Gaps	7
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Db)	59	PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP	118
Żу	•	116	APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKR	166
Db)	119	APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 1	178
Qу		167	RGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSL 2	226
Db)	179	RGSS	182
Qу		227	SPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYS 2	286
Db)	183		182
Qу		287	EMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEK	346
Db	ı	183		182
Qу		347	TMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLED 4	106
Db	1	183		182
Qу		407	SLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTS 4	166
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Qу		467	ENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEG 5	526
Db		183		182
Qу		527	LTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSP 5	586
Db		183		182
Qу		587	VLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALK 6	546
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Qу
        183 ----- 182
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        887 ELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPOTEMGSIVKSKSLTK 946
QУ
        183 ------ 185
Db
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                                     Db
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Qу
       1007 IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSA 1066
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        217 IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVOKYSNSA 276
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            Db
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       1127 ERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
Ov
            337 ERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 373
RESULT 12
US-10-060-036-72
; Sequence 72, Application US/10060036
; Publication No. US20030073144A1
; GENERAL INFORMATION:
; APPLICANT: Benson, Darin R.
 APPLICANT: Kalos, Michael D.
 APPLICANT: Lodes, Michael J.
 APPLICANT: Persing, David H.
  APPLICANT: Hepler, William T.
  APPLICANT: Jiang, Yuqiu
  TITLE OF INVENTION: COMPOSITIONS AND METHODS FOR THE THERAPY
  TITLE OF INVENTION: AND DIAGNOSIS OF PANCREATIC CANCER
  FILE REFERENCE: 210121.566
  CURRENT APPLICATION NUMBER: US/10/060,036
  CURRENT FILING DATE: 2002-01-30
  NUMBER OF SEQ ID NOS: 4560
  SOFTWARE: FastSEO for Windows Version 4.0
; SEQ ID NO 72
   LENGTH: 373
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US-10-060-036-72
 Query Match 20.4%; Score 1191; DB 15; Length 373; Best Local Similarity 27.8%; Pred. No. 1.4e-52;
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Matches 327; Conservative 12; Mismatches 20; Indels 818; Gaps

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Db	59	PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPERQPSWDPSPVSSTVP	118
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Db	183		182
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Db	183		182
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                                       Db
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             Db
         217 IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSA 276
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Qу
             277 LGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSVPVIY 336
Db
Qу
        1127 ERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
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         337 ERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 373
RESULT 13
US-09-893-348-21
; Sequence 21, Application US/09893348
; Patent No. US20020072493A1
; GENERAL INFORMATION:
  APPLICANT: EISENBACH-SCHWARTZ, Michal
  APPLICANT: COHEN, Irun R.
             BESERMAN, Pierre
  APPLICANT:
  APPLICANT: MOSONEGO, Alon
  APPLICANT:
             MOALEM, Gila
  TITLE OF INVENTION: ACTIVATED T-CELLS, NERVOUS SYSTEM-SPECIFIC ANTIGENS AND
  FILE REFERENCE: EIS-SCHWARTZ=2A
  CURRENT APPLICATION NUMBER: US/09/893,348
  CURRENT FILING DATE: 2001-06-28
  PRIOR APPLICATION NUMBER: US 09/314,161
  PRIOR FILING DATE: 1999-05-19
  PRIOR APPLICATION NUMBER: US 09/218,277
  PRIOR FILING DATE: 1998-12-22
  PRIOR APPLICATION NUMBER: PCT/US98/14715
  PRIOR FILING DATE: 1998-07-21
  PRIOR APPLICATION NUMBER: IL 124500
  PRIOR FILING DATE: 1998-05-19
  NUMBER OF SEQ ID NOS: 29
  SOFTWARE: PatentIn version 3.1
 SEQ ID NO 21
   LENGTH: 199
   TYPE: PRT
   ORGANISM: Rattus norvegicus
US-09-893-348-21
 Query Match
                       15.8%;
                              Score 925; DB 9; Length 199;
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99.0%; Pred. No. 1.7e-39;

Best Local Similarity

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Matches 189; Conservative
                             0; Mismatches
                                             2; Indels
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Qу
               9 KDKVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVI 68
Db
        1033 QAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKF 1092
Qу
             Db
          69 QAIQKSDEGHPFRAYLESEVAISEELVOKYSNSALGHVNSTIKELRRLFLVDDLVDSLKF 128
Qу
        1093 AVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIO 1152
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         189 AKIPGLKRKAD 199
RESULT 14
US-09-893-348-25
; Sequence 25, Application US/09893348
; Patent No. US20020072493A1
; GENERAL INFORMATION:
  APPLICANT: EISENBACH-SCHWARTZ, Michal
  APPLICANT: COHEN, Irun R.
  APPLICANT: BESERMAN, Pierre
  APPLICANT: MOSONEGO, Alon
  APPLICANT: MOALEM, Gila
  TITLE OF INVENTION: ACTIVATED T-CELLS, NERVOUS SYSTEM-SPECIFIC ANTIGENS AND
THEIR USES
  FILE REFERENCE: EIS-SCHWARTZ=2A
  CURRENT APPLICATION NUMBER: US/09/893,348
  CURRENT FILING DATE: 2001-06-28
  PRIOR APPLICATION NUMBER: US 09/314,161
  PRIOR FILING DATE: 1999-05-19
  PRIOR APPLICATION NUMBER: US 09/218,277
  PRIOR FILING DATE: 1998-12-22
  PRIOR APPLICATION NUMBER: PCT/US98/14715
  PRIOR FILING DATE: 1998-07-21
  PRIOR APPLICATION NUMBER: IL 124500
  PRIOR FILING DATE: 1998-05-19
  NUMBER OF SEQ ID NOS: 29
  SOFTWARE: PatentIn version 3.1
 SEO ID NO 25
   LENGTH: 199
   TYPE: PRT
   ORGANISM: Homo sapiens
US-09-893-348-25
 Query Match
                       15.5%; Score 908; DB 9; Length 199;
 Best Local Similarity
                      96.3%; Pred. No. 1.2e-38;
 Matches 184; Conservative
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Db
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1033 QAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKF 1092
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        1093 AVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHOVOIDHYLGLANKSVKDAMAKIO 1152
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Db
        1153 AKIPGLKRKAD 1163
Qу
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Db
         189 AKIPGLKRKAE 199
RESULT 15
US-09-789-386-4
; Sequence 4, Application US/09789386
; Patent No. US20020010324A1
; GENERAL INFORMATION:
  APPLICANT: MICHALOVICH, DAVID
  APPLICANT: PRINJHA, RABINDER KUMAR
  TITLE OF INVENTION: NOVEL COMPOUNDS
  FILE REFERENCE: GP-30165-C1
  CURRENT APPLICATION NUMBER: US/09/789,386
  CURRENT FILING DATE: 2001-02-21
  PRIOR APPLICATION NUMBER: U.K. 9916898.1
  PRIOR FILING DATE: 1999-07-19
  PRIOR APPLICATION NUMBER: U.K. 9816024.5
  PRIOR FILING DATE: 1998-07-22
  PRIOR APPLICATION NUMBER: US 09/359,208
  PRIOR FILING DATE: 1999-07-22
  NUMBER OF SEO ID NOS: 6
  SOFTWARE: FastSEQ for Windows Version 3.0
 SEO ID NO 4
   LENGTH: 289
   TYPE: PRT
   ORGANISM: HOMO SAPIENS
   FEATURE:
   NAME/KEY: UNSURE
   LOCATION: (31)(138)
US-09-789-386-4
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                       15.2%; Score 888; DB 9; Length 289;
 Best Local Similarity 63.9%; Pred. No. 2e-37;
 Matches 184; Conservative 34; Mismatches 58;
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            Db
          3 MDLKEQPGNTISAGQEDFPSVLLETAASXPSLSPLSAASFKEHEYLGNLSTVLPTEGTLO 62
         255 ETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVR 314
Qу
            Db
         63 ENVSEASKEVSEKAKTLLIDRDLTEFSELEYSEMGSSFSVSPKAESAVIVANPREEIIVK 122
         315 SKDKED-LVCSAALHSPQESPVG----KEDRVVSPEKTMDIFNEMQMSVVAPVREEYAD 368
Qу
            :||:|: || : || || ||
                                     111 111 11 1 11 :::| ||:|||||
         123 NKDEEEKLVSNNILHXQQELPTALTKLVKEDEVVSSEKAKDSFNEKRVAVEAPMREEYAD 182
Db
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QУ		FKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDA 424
Db	183	FKPFERVWEVKDSKEDS-DMLAAGGKIESNLESKVDKKCFADSLEQTNHEKDSESSNDDT 241
QУ	425	SFPSTPEPVKDSSRAYITCASFT-SATESTTANTFPLLEDHTSENKTD 471
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Search completed: December 19, 2003, 15:42:41

Job time : 45 secs

GenCore version 5.1.6 Copyright (c) 1993 - 2003 Compugen Ltd.

OM protein - protein search, using sw model

Run on:

December 19, 2003, 15:33:28; Search time 27 Seconds

(without alignments)

4142.377 Million cell updates/sec

Title:

US-09-830-972-2

Perfect score: 5848

Sequence:

1 MEDIDQSSLVSSSTDSPPRP.....VKDAMAKIQAKIPGLKRKAD 1163

Scoring table:

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Gapop 10.0 , Gapext 0.5

Searched:

283308 seqs, 96168682 residues

Total number of hits satisfying chosen parameters: 283308

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

PIR_76:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

왕

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		۰				· ·	
Result		Query					
No.	Score	Match	Length	DB	ID	_	Description
1	789.5	-12 -	776		7.46502		
_	,	13.5	776	2	A46583		neuroendocrine-spe
2	685	11.7	208	2	I6090 4		neuroendocrine-spe
3	671	11.5	267	2	A60021		tropomyosin-relate
4	517	8.8	2484	2	T26216		hypothetical prote
5	503.5	8.6	2607	2	T26215		hypothetical prote
6	328.5	5.6	5327	2	T13564		microtubule-associ
7	322	5.5	7962	2	I38346		elastic titin - hu
8	320	5.5	222	2	T26213		hypothetical prote
9	304.5	5.2	873	2	A47283		calphotin - fruit
10	302.5	5.2	1829	2	T24583		hypothetical prote
11	295.5	5.1	865	2	A47282	•	calcium-binding pr
12	292	5.0	3507	2	T34513		hypothetical prote
13	291.5	5.0	2364	2	A56577		microtubule-associ

14	288.5	4.9	971	2	T19431
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17	279.5	4.8	1262	2	T22523
18	277	4.7	1621	2	A82255
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23	273	4.7	1558	2	B71603
24	272.5	4.7	3534	2	T42567
25	270	4.6	1230	2	T22458
26	269.5	4.6	2187	2	T30826
27	267	4.6	1684	2	JW0057
28	265.5	4.5	1828	2	A40115
29	263	4.5	1825	2	S13507
30	261.5	4.5	1087	1	QFMSH
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36	258.5	4.4	1830	2	A37981
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38	256	4.4	2361	2	T25752
39	254.5	4.4	6642	2	T29757
40	254	4.3	1616	2	G64242
41	254	4.3	3381	2	T42389
42	253	4.3	3421	1	WZBEB6
43	252.5	4.3	5170	2	T15348
44	251	4.3	1824	1	QRHUMT
45	250.5	4.3	4377	2	A55575

hypothetical prote microtubule-associ hypothetical prote hypothetical prote hypothetical prote ankyrin 2, neurona hypothetical prote mucin-like protein hypothetical prote RESA-H3 antigen PF tegument protein 2 hypothetical prote nascent polypeptid gravin - human microtubule-associ microtubule-associ neurofilament trip neurofilament trip fimbriae-associate nestin - golden ha neurofilament trip hypothetical prote microtubule-associ microtubule-associ hypothetical prote protein UNC-89 - C cytadherence-acces versican precursor 367K tegument prot hypothetical prote microtubule-associ ankyrin 3, long sp

ALIGNMENTS

RESULT 1 A46583

neuroendocrine-specific protein, splice form A - human

N; Contains: neuroendocrine-specific protein, splice form B

C; Species: Homo sapiens (man)

C;Date: 24-May-1996 #sequence_revision 24-May-1996 #text_change 05-Nov-1999

C; Accession: A46583; I60903

R;Roebroek, A.J.; van de Velde, H.J.; Van Bokhoven, A.; Broers, J.L.; Ramaekers, F.C.; Van de Ven, W.J.

J. Biol. Chem. 268, 13439-13447, 1993

A; Title: Cloning and expression of alternative transcripts of a novel

neuroendocrine-specific gene and identification of its 135-kDa translational product.

A; Reference number: A46583; MUID: 93293865; PMID: 7685762

A; Accession: A46583

A; Status: preliminary; translated from GB/EMBL/DDBJ

A; Molecule type: mRNA A; Residues: 1-776 < ROE1>

A; Cross-references: GB:L10333; NID:g307306; PIDN:AAA59950.1; PID:g307307

A; Accession: I60903

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A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 421-776 < ROE2>
A; Cross-references: GB:L10334; NID:q307308; PIDN:AAA59951.1; PID:q307309
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A; Gene: GDB: RTN1; NSP
A; Cross-references: GDB:203968; OMIM:600865
A; Map position: 14q21-14q22
  Query Match
                     13.5%; Score 789.5; DB 2; Length 776;
 Best Local Similarity 31.2%; Pred. No. 3.4e-25;
 Matches 243; Conservative 115; Mismatches 252; Indels 169; Gaps
        487 TSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGT 546
QУ
           65 SGPARQSP--VAMETASTGVAGVSSAMDHTFSTTSKDGEG-----SCYTSLI----S 110
Db
Qу
        547 KIAYETKVDLVQTSEAIQ-ESLYPTAQLCPSFEEAEATPSPVLPDI--VMEAPLNS---- 599
            111 DICYPPQEDSTYFTGILQKENGHVTISESP---EELGTPGPSLPDVPGIESRGLFSSDSG 167
Db
Qу
        600 --LLPSAGASVVQPSVSPLEAPPPVSY-----DSIKLEPENPPPYEEA------------ 641
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        168 IEMTPAESTEVNKILADPLDOMKAEAYKYIDITRPEEVKHQEQHHPELEDKDLDFKNKDT 227
        642 NVALKALGTKEGIKEPE-----SFNAAVQETEAPYISIACDLIKETKLSTE-PSP 690
Qу
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                                  228 DISIK----PEGVREPDKPAPVEGKIIKDHLLEESTFAPYID---DLSEEQRRAPQITTP 280
Db
        691 DFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLF-SDDSIPEV----PQTQEEAVMLMK 745
Qу
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Db
        281 VKITLTEIE-----PSVETTTQEKTPEKQDICLKPSPDTVPTVTVSEPEDDSPGSITPP 334
Qу
        746 ESLTEVSETVAQHK----EERLSASPQELGKPYLESFOP----NLHSTKDAASND---- 792
            Db
        335 SSGTEPSAAESQGKGSISEDELITAIKEAKGLSYETAENPRPVGQLADRPEVKARSGPPT 394
Qу
        793 IPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKI----- 827
           395 IPSPLDHEASSAESGDSEIELVSEDPM--AAEDALPSGYVSFGHVGGPPPSPASPSIQYS 452
Db
        828 ---KESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEI----ANIOSG 879
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             453 ILREEREAELDS---ELIIESCDASSASEESPKREQDSPPMKPSALDAIREETGVRAEER 509
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        880 ADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIV 939
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              510 APSRRGLAEPG--SFLD-YPSTEPQPGPEL------PPGDGALEPETPM--- 549
Db
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Qу
                     1:| :|||||
Db
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        985 IKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPF 1044
QУ
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        1105 LFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
Qу
            :
                                                  718 LFNGLTLLLMAVVSMFTLPVVYVKHQAQIDQYLGLVRTHINAVVAKIQAKIPGAKRHAE 776
Db
RESULT 2
I60904
neuroendocrine-specific protein C - human
C; Species: Homo sapiens (man)
C; Date: 24-May-1996 #sequence revision 24-May-1996 #text change 05-Nov-1999
C; Accession: I60904
R; Roebroek, A.J.; van de Velde, H.J.; Van Bokhoven, A.; Broers, J.L.; Ramaekers,
F.C.; Van de Ven, W.J.
J. Biol. Chem. 268, 13439-13447, 1993
A; Title: Cloning and expression of alternative transcripts of a novel
neuroendocrine-specific gene and identification of its 135-kDa translational
A; Reference number: A46583; MUID: 93293865; PMID: 7685762
A; Accession: I60904
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 1-208 < RES>
A; Cross-references: GB:L10335; NID:g307310; PIDN:AAA59952.1; PID:g307311
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A; Gene: GDB: RTN1; NSP
A; Cross-references: GDB:203968; OMIM:600865
A; Map position: 14q21-14q22
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QУ
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            Db
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        1093 AVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQ 1152
Qу
            138 AVLMWLLTYVGALFNGLTLLLMAVVSMFTLPVVYVKHQAQIDQYLGLVRTHINAVVAKIQ 197
Db
        1153 AKIPGLKRKAD 1163
Qу
            |||||:
        198 AKIPGAKRHAE 208
Dh
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RESULT 3 A60021

tropomyosin-related protein, neuronal - rat C;Species: Rattus norvegicus (Norway rat)

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C; Accession: A60021
R; Wieczorek, D.F.; Hughes, S.R.
Brain Res. Mol. Brain Res. 10, 33-41, 1991
A; Title: Developmentally regulated cDNA expressed exclusively in neural tissue.
A; Reference number: A60021; MUID: 91278684; PMID: 1647480
A; Accession: A60021
A; Molecule type: mRNA
A; Residues: 1-267 <WIE>
A;Cross-references: EMBL:X52817; NID:g456549; PIDN:CAA37001.1; PID:g456550
C; Comment: This neuronal-specific mRNA was identified by hybridization to an
alpha-tropomyosin probe but does not show homology in amino acid sequence.
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  Best Local Similarity
                         66.3%; Pred. No. 5.4e-21;
  Matches 124; Conservative 33; Mismatches 30; Indels
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Qу
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QУ
              -:||;|:||||||
Db
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             Db
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        1153 AKIPGLK 1159
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Db
         189 AKIPGAR 195
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T26216
hypothetical protein W06A7.3c - Caenorhabditis elegans
C; Species: Caenorhabditis elegans
C;Date: 15-Oct-1999 #sequence revision 15-Oct-1999 #text change 15-Oct-1999
C; Accession: T26216
R; Ainscough, R.
submitted to the EMBL Data Library, August 1996
A; Reference number: Z20173
A; Accession: T26216
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-2484 <WIL>
A; Cross-references: EMBL: Z78066; PIDN: CAB51467.1; GSPDB: GN00023; CESP: W06A7.3c
A; Experimental source: clone W06A7
C; Genetics:
A; Gene: CESP: W06A7.3c
A; Map position: 5
A; Introns: 1827/1; 1866/3; 1963/3; 1990/3; 2262/1; 2289/1; 2339/2; 2463/2
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 Best Local Similarity
                        20.1%; Pred. No. 2.6e-13;
 Matches 262; Conservative 189; Mismatches 413; Indels 440; Gaps
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C;Date: 03-Mar-1993 #sequence revision 03-Mar-1993 #text change 05-Nov-1999

Qy	28	FVTEPEDEEDEEEEEDEEEDDEDLEELEVL	57
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Db	1497	DASDVNEQDEESTLKILKVVPSEPSLLELDFTNDPKVIHVPIPLMEPATMYLEEMVEW	1554
Qу	109	ERSPAAPAPSLPPAAAVLPSKLP-EDDEPPARPPPPPPA	146
Db	1555	: : :	1614
Qy	147	GASPLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPS-SAEKIMDLMEQPGNTV	205
Db	1615	: : :	1670
Qу	206	SSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELP	265
Db	1671	:	1695
Qy	266	ERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKE	319
Db	1696	EKPTEDIGALSPLSPNTLAEYEEVPMMDMQS	1726
Qy	320	DLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQ-MSVVAPVREEYADFKPFEQAWEV	378
Db	1727	VPHSPQEKQEEIEALSEIIEEPQAMKEVEKPVESAPE-	1763
Qу	379	KDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDS	436
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Qy	437	SRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFL : :	496
Db	1802	DDDGSECLDSIGDLSERTIQRFN	1824
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Db	1825	TSIDDPSIRRDSFSSISSFGDRQKFRTAIENIRQDLLP	1862
Qy	549	AYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASV :::	608
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Qу	609	VQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAA	663
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Qу	664	VQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPE: :::: ::: :	707 .
Db	1938	LDEFAAPPVHDPMQKSVFGSLGSDDMKPGSQDDGFVFIERNEANEATLKKNQKMSSH	1994
Qу	708	HAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTE	750
Db	1995		2054

```
Qу
        751 VSETVAOHKE-----ERLSASPQELGKPYLESFQP 780
            | : : | | |
                                             | | : : | : |
        2055 VEQKIEQVKEPIVDSLHKAYDGVGDFVHETVPNAVDDFVREAEKQLPESPVPEKIETPEP 2114
Db
        781 --NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDK-----IKES 830
Qу
              ::| | | ::: ::|
        2115 LVDIHDTVDKVHDEVDNFLRREP-----TPPFETDDVAPLSDDKPQFGNQTPEEDE 2165
Db
        831 ETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGA----DSLPCL 886
Qу
             2166 TTFDRKGPLTIPEEVEKAAAAQNND---LDDFDPLVTSNTGAAFGAAVGAAAAVESLTEE 2222
Db
        887 ELPCDLSFKNI-----YPKDEVHVSDEFSENRSSVSKASISPSNVSA-----LEPOTEM 935
Qу
                          : || |:
                 1: :
        2223 EMFGHQKFETVPRPPTPPKD---ISDE-----DVKPSTVNLGPSHHHSHPSSPH 2268
Db
        936 GSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGAS 995
Qу
                                    Db
       2269 HSILKHHG-----
                            ------DAWIDFKTVPPCVLDVIYWRDAKKSAIVLSLA 2308
        996 LFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAIS 1055
Qу
            Db
       2309 LLVLFVLAKYPLLTVVTYSLLLALGAAAGFRVFKKVEAQIKKTDSEHPFSEILAQDLTLP 2368
       1056 EELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILA 1115
Ov
            Db
       2369 QEKVHAQADVFVEHATCIANKLKKLVFVESPLESIKFGLVLWSLTYIASWFSGFTLAILG 2428
       1116 LISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLK 1159
Qу
            Db
       2429 LLGVFSVPKVYESNQEAIDPHLATISGHLKNVQNIIDEKLPFLR 2472
RESULT 5
T26215
hypothetical protein W06A7.3a - Caenorhabditis elegans
C; Species: Caenorhabditis elegans
C;Date: 15-Oct-1999 #sequence revision 15-Oct-1999 #text change 15-Oct-1999
C; Accession: T26215
R; Ainscough, R.
submitted to the EMBL Data Library, August 1996
A; Reference number: Z20173
A; Accession: T26215
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-2607 <WIL>
A;Cross-references: EMBL:Z78066; PIDN:CAB01522.2; GSPDB:GN00023; CESP:W06A7.3a
A; Experimental source: clone W06A7
C; Genetics:
A; Gene: CESP: W06A7.3a
A; Map position: 5
A;Introns: 1827/1; 1866/3; 1963/3; 1990/3; 2262/1; 2289/1; 2412/1; 2462/2;
2586/2
 Query Match
                       8.6%; Score 503.5; DB 2; Length 2607;
 Best Local Similarity 20.8%; Pred. No. 1e-12;
 Matches 280; Conservative 181; Mismatches 481; Indels 407; Gaps 52;
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Qy	28	FVTEPEDEEDEEEEEDEEEDDEDLEELEVL	57
Db	1437	FGTESSEESQKADGNQENQEEEDVVAELNFHPIRQWRDEDVISLQSLKSLVAEVGCITDV	1496
Qу	58	ERKPAAGLSAAAVPPAAAAPL-LDFSSDSVPPAPRGPLPAAPPAAPERQPSW	108
Db	1497	DASDVNEQDEESTLKILKVVPSEPSLLELDFTNDPKVIHVPIPLMEPATMYLEEMVEW	1554
Qу	109	ERSPAAPAPSLPPAAAVLPSKLP-EDDEPPARPPPPPPA	146
Db	1555	IIADAVKEVSEMEVVTESEISEMAPQVSESTCPIPEPLADLKLPVEDDEKTPEPEPVVPG	1614
Qy	147	GASPLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPS-SAEKIMDLMEQPGNTV	205
Db	1615	QVQERIIPIEVEQAPTIPQRPPRAPKSELPKVAKPLDDSKSRVRFAPLNIKLGRTY	1670
Qy	206	SSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELP	265
Db	1671	SEEQQKELVESLERPLTIITQQKPP	1695
Qy	266	ERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKE	319
Db	1696	:	1746
Qy .	320	DLVCSAALHSPQES-PVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQA : : : : : : : :	375
Db	1747	: : : : : : : : EPQAMKEVEKPVESAPEKDNESLEAPEIINEPIRRVLVETKIMGPGKSL	1795
Qу	376	WEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSE	418
Db	1796	NEDNDDDDDGSECLDSIGDLSERTIQR-FNTSIDDPSIRRDSFSSISSFGDRQKFRTAIE	1854
QУ	419	GRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLE	462
Db	1855	: : : : : : :	1911
Qy	463	TEKTSPKTSNPF	495
Db	1912	: : :: : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : : :	1971
Qy	496	LVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQ	533
Db	1972	:: : : : : : : : :	2031
Qy	534	EACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVL	588
Db	2032	: : :: : : : : SASEYKKQAVDSGDEIGRELLDNVEQKIEQVKEPIVDSLHKAYDGVGDFVH-ETVPNAV-	2089
Qу	589	PDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKAL	648
Db	2090		2106
Qy	649	GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEK	703
Db	2107	: : : : : EKIETPEPLVDIHDTVDKVHDEVDNFLRREPTPPFE	2142

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Qу
         704 SVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLT---EVSETVAQHKE 760
                     : | : | | : : | | : : | | | | | | : |
Db
        2143 -----TDDVAPLSDDKPQFGNQT-PE----EDETTFDRKGPLTIPEEVEKAAAAQNN 2189
Qу
         761 ERLSASPQELGKPYLESFQPNLHSTKDA-----ASNDIPTLTKKEKISLQMEEFNTAI 813
                        Db
        2190 D-----LDDFDPLVTSNTGAAFGAAVGAAAAVESLTEEEMFGHQ--KFETVP 2234
         814 YSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDSP--KLAKEYTDLEVSDKS 871
Qу
                                     :: |: : |: : |:
        2235 RPPTPPKDISDEDVKPS-----TVNLGPSHHHSHPSSPHHSILKHHGDAWIDFKT 2284
Db
         872 EIANIQSG------ADSLPCLE----LPCDLSFKNIYPKDEVH 904
Qу
                                        : | | | :
                                                      Db
        2285 VPPCAQNAFSPGEIMFLLAFFVYLSCFASFFSKSLPLLDNLLSLVVYLSISLI----IH 2339
         905 VSDE-----FSENRSSVSKASISPSNVSA-----LEPOTEMGSIVKSKSLTKEAEK 950
QУ
                       | :::||
                                   :
                                                  : :|
        2340 VKHHRKFRWNEEQATTMSKLGAVGRGLYALIAFIVNIVLRVGLNVALVVGVAVSAHEAYK 2399
Db
         951 KLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSV 1010
QУ
                           -----SGVLRKKEVLDVIYWRDAKKSAIVLSLALLVLFVLAKYPLLTV 2446
Db
        2400 LTKS-
        1011 TAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHV 1070
Qу
              2447 VTYSLLIALGAAAGFRVFKKVEAQIKKTDSEHPFSEILAQDLTLPQEKVHAQADVFVEHA 2506
Db
        1071 NSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQ 1130
QУ
                 Db
        2507 TCIANKLKKLVFVESPLESIKFGLVLWSLTYIASWFSGFTLAILGLLGVFSVPKVYESNQ 2566
        1131 VQIDHYLGLANKSVKDAMAKIQAKIPGLK 1159
Qy
              Db
        2567 EAIDPHLATISGHLKNVONIIDEKLPFLR 2595
RESULT 6
T13564
microtubule-associated protein homolog - fruit fly (Drosophila melanogaster)
N; Alternate names: hypothetical protein EG: 49E4.1
C; Species: Drosophila melanogaster
C;Date: 13-Aug-1999 #sequence_revision 13-Aug-1999 #text change 17-Nov-2000
C; Accession: T13564
R; Spanos, L.; Papagiannakis, G.; Siden-Kiamos, I.; Louis, C.
submitted to the EMBL Data Library, April 1999
A; Description: Sequencing the distal X chromosome of Drosophila melanogaster.
A; Reference number: Z17689
A; Accession: T13564
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-5327 <SPA>
A; Cross-references: EMBL: AL031128; PIDN: CAA20006.1
C; Genetics:
A; Cross-references: FlyBase: FBgn0025392
A; Introns: 24/2; 52/3; 104/3; 179/1; 232/1; 1669/3; 2566/1; 4798/3; 5272/1
A; Note: EG: 49E4.1
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C;S	uperfamily	: Drosophila 576K microtubule-associated protein homolog	
В	est Local :	5.6%; Score 328.5; DB 2; Length 5327; Similarity 22.7%; Pred. No. 4.1e-05; 5; Conservative 176; Mismatches 453; Indels 238; Gaps	54;
Qу		EDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVL	·
Db			
		DEADKSKEESRRESGAEKSPLASKEASRPASVAESIKDEAEKSKEESRRESVAEKSPL	
Qу		ERKPAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAP	
Db	3254	PSKEASRPTSVAKSVKDEAEKSKEESSRDSVAEKSPLASKEASRPASVAESVQDEAEK	3311
Qу	103	ERQPSWERSPAAPAPSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLA-EPAA	156
Db	3312	SKEESRRESVAEKSPLAYKEASRP-ASVAESIKDEAEKSKEESRRESVAEKSPLASKEAS	3370
Qy	157	PPSTPAAPKRRGSGSVDETLFALPAAS-EPVIPSS-AEKIMDLMEQPGNTVSSG	208
Db	3371	RPTSVAESVKDEAEKSKEESSRDSVAEKSPLASKEASRPASVAESVQDEAEKSKEESRRE	3430
Qу	209	QEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEAS	261
Db	3431	: : : : SVAEKSPLASKEASRPASVAESVKDDAEKSKEESRRESVAEKSPLASKEASRPASVAESV	3490
Qу	262	KELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKE	319
Db	3491	: :: : : : : : : : :	3541
Qу	320	DLVCSAALHSPQES-PVGKEDRVV-SPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWE	377
Db	3542	: :: : :	3600
Qу	378	VKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNE	422
Db	3601	:: : : : : : : VKDEAEKSKEVSRRESVAEKSPLPSKEASRPTSVAESVKDEADKSKEESRRESGAEKSPL	3660
Qу	423	DASFP-STPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTD-	471
Db	3661	:	3715
Qу	472	-EKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEG-	526
Db	3716	: : : : ::	3769
Qу		LTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYP	
Db	3770	: : : : : : KAESIKGDQSSLASKETSRPDSVVESVKDETEKPEGSAIDKSQVASRPESVAVSAKD	3826
Qу	570	TAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKL	629
Db	3827	: : : :	3872
Qy	630	EPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPS	689
Db	3873	:::	3911

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690 PDFSNYSEIAKFEKSVPEH--AELVEDSSPESEPVDLF-----SDDSIPEV 733
Qу
            3912 P--AVLSKPAEFSQPDTGHTASTPVDEASPVLEEIEVVEQHTTSGVGATGATAETDLLDL 3969
Db
Qу
        734 PQTQEEAVMLMKESLTEVSETVAQHKEERLSASPOELGKPYLESFOPNLHSTKDAASNDI 793
            3970 TETKSETV--TKQSETTLFETLTSKVESKVEVLESSV-KQVEEKVQTSVKQAETTVTDSL 4026
Db
        794 PTLTKKEKISLQMEE-----FNTAIYSNDDLLSSKEDKIKESETFSDSSP 838
QУ
                                    Db
       4027 EQLTKKS--SEQLTEIKSVLDTNISNVTNLFSTAVETIEKKVQDVTEKVIEKAT----- 4078
        839 IEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIY 898
QУ
            4079 - EHVSEHVT--TTGESSTETSQEKSSLDLGTFSELRETHITTVGSPEFTV-----TIC 4128
Db
Qу
        899 PKDE--VH-VSDEFSENR----SSVSKAS-ISPSNVSALEPQTE-----MGSIVKS- 941
            : ::[]
       4129 ERDEPVLHDIKEEDEEHRFSPPSDVDKAAIIPPQPMRPLSPREEEVAKIVADVAKVLKSD 4188
Db
        942 KSLT-----KEAEKKLPS--DTEKE-DRSLSAVLSAELS 972
Qу
            Db
       4189 KDITDIIPDFDERQLEEKLKSTADTEEESDKSTRDEKSLEIS 4230
RESULT 7
I38346
elastic titin - human (fragment)
C; Species: Homo sapiens (man)
C;Date: 29-May-1998 #sequence_revision 29-May-1998 #text change 21-Jul-2000
C; Accession: I38346
R; Labeit, S.; Kolmerer, B.
Science 270, 293-296, 1995
A; Title: Titins: giant proteins in charge of muscle ultrastructure and
A; Reference number: A57430; MUID: 96026330; PMID: 7569978
A; Accession: I38346
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: mRNA
A; Residues: 1-7962 < RES>
A;Cross-references: EMBL:X90569; NID:g1017426; PIDN:CAA62189.1; PID:g1017427
C; Genetics:
A; Gene: GDB: TTN
A; Cross-references: GDB:127867; OMIM:188840
A; Map position: 2g31-2g31
 Query Match
                      5.5%; Score 322; DB 2; Length 7962;
 Best Local Similarity 21.7%; Pred. No. 0.00013;
 Matches 259; Conservative 146; Mismatches 416; Indels 370; Gaps
         23 AFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVL-----ERKPAAGL, 65
Qу
           6574 AFEEEVVTHVEEYLVEEEEEYIHEEEEFITEEEVVPVIPVKVPEVPRKPVPEEKKPVPVP 6633
Db
         66 SAAAVPPAAA-----APLLDFSSDSVPPA----PRGPLP-----AAP 98
Qу
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	Db	6634	KKKEAPPAKVPEVPKKPEEKVPVLIPKKEKPPPAKVPEVPKKPVPEEKVPVPVPKKVEAP	6693
	Qy	99	PA-APERQPSWERSPAAPAPSLPPA-AAVLPSKL-PEDDEPPARPPPPP	144
	Db	6694	:: : : : : : PAKVPEVPKKPVPEKKVPVPAPKKVEAPPAKVPEVPKKLIPEEKKPTPVPKKVEAPPPKV	6753
	Qy	145	PAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNT	204
	Db	6754	PKKREPVPVPVALPQEEEVLFEEEIVPE-EEVLPEEEE	6790
	Qy	205	VSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNE : :	259
	Db	6791	VLPEEEEVLPEEEEVLPEEEEIPPEEEEVPPEEEYVPEEEEFVPEEEVLPE	6841
Q	Qy	260	ASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIV :: : : : : : : : : : : : : : :	313
	Db	6842	VKPKVPVPAPVPEIKKKVTEKKVVIPKKEEAPPAKVPEVPKKVEEKRII	6890
	Qy	314	RSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEM	354
	Db	6891	LPKEEEVLPVEVT-EEPEEEPISEEEIPEEPPSIEEVEEVAPPRVPEVIKKAVPEAPTPV	6949
	Qy	355	QMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESK :	398
	Db	6950	PKKVEAPPAKVSKKI PEEKVPVPVQKKEAPPAKVPEVPKKVPEKKVLVPKKEAVPPA	7006
	Qy	399	VDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCA	444
	Db	7007	KGRTVLEEKVSVAFRQEVVVKERLELEVVEAEVEEIPEEEEFHEVEEYFEEG	7058
	Qy	445	SFTSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKT	491
	Db	7059	EFHEVEEFIKLEQHRVEEEHRVEKVHRVIEVFEAEEVEVFEKPKAPPKGPEISEKIIPPK	7118
	Qy	492	SNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLV	532
	Db	7119	KPPTKVVPRKEPPAKVPEVPKKIVVEEKVRVPEEPRVPPTKVPEVLPPKEVVPEKKVPVP	7178
	Qy	533	QEACESELNEATGTKIAYETKVDL	569
	Db	7179	PAKKPEAPPPKVPEAP-KEVVPEKKVPVPPPKKPEVPPTKVPEVPKAAVPEKKVPEAIPP	7237
	Qy	570	TAQLCPSFEEAEATPSPV-LPDIVMEAPLNSLLPSAGASVVQPSVSPLE : : : : : : : :	617
	Db	7238	KPESPPPEVFEEPEESPSAPPKKPEVPPVRVPEVPKEVVPEKKVPAAPPKKPEVTPVK	7295
	QУ	618	APDSIKLEPENPPPYEE	639
	Db	7296	VPEAPKEVVPEKKVPVPPPKKPEVPPTKVPEVPKVAVPEKKVPEAIPPKPESPPPEVFEE	7355
	Qу		AMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEP	
	Db		PEEVALEE-PPAEVVEEPEPAAPPQVTVPPKNPVPEKKAPAVVAKKPELPPVK	
	Qу	689	SPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESL : : : : : : : : : : : : : : : : : : :	748
	Db	7408	VPEVP - KEVVP - EKKVP I VVPKKPEAPPA KVPEVP KEVV	7444

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749 TEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEE 808
Qу
                : |:|: : |
Db
       7445 PEKKVAVPKKPEVPPAKVPEVPKKPVLEE-KPAVPVPERAESPPPEVYEEPEEIAPEEE- 7502
        809 FNTAIYSNDDLLSSKEDK---IKESE-----TFSDSSPIEIIDEFPTFVSAKDDSPKLA 859
Oy
                    Db
       7503 -----IAPEEEKPVPVAEEEEPEVPPPAVPEEPKKIIPEKKVPVIKKPEAPP-P 7550
        860 KEYTDLEVSDKSEIANIOSGADSLPCLELPCDLSFK-NIYPKDEVHVSDEFSENRSSVSK 918
QУ
           7551 KEPEPEKVIEKPKLKPRPPPPPPAPPKEDVKEKIFQLKAIPKKKV-----PENPQVPEK 7604
Db
        919 ASISPSNVSALEPOTEMGSIVKSKSLTKEAEKKLP--SDTEKEDRSLSAVL 967
Qу
             7605 VELTPLKVPGGE-----KKVRKLLPERKPEPKEEVVLKSVL 7640
Dh
RESULT 8
T26213
hypothetical protein W06A7.3b - Caenorhabditis elegans
C; Species: Caenorhabditis elegans
C; Date: 15-Oct-1999 #sequence revision 15-Oct-1999 #text change 15-Oct-1999
C:Accession: T26213
R; Ainscough, R.
submitted to the EMBL Data Library, August 1996
A; Reference number: Z20173
A; Accession: T26213
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-222 <WIL>
A; Cross-references: EMBL: Z78066; PIDN: CAB01523.1; GSPDB: GN00023; CESP: W06A7.3b
A; Experimental source: clone W06A7
C; Genetics:
A; Gene: CESP: W06A7.3b
A; Map position: 5
A; Introns: 27/1; 77/2; 201/2
                     5.5%; Score 320; DB 2; Length 222;
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 Best Local Similarity 32.1%; Pred. No. 1e-06;
 Matches 59; Conservative 47; Mismatches 78; Indels 0; Gaps
        976 VVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAI 1035
Qу
           Db
         27 ILDVIYWRDAKKSAIVLSLALLVLFVLAKYPLLTVVTYSLLLALGAAAGFRVFKKVEAOI 86
Qу
       1036 QKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVL 1095
           Db
         87 KKTDSEHPFSEILAODLTLPOEKVHAOADVFVEHATCIANKLKKLVFVESPLESIKFGLV 146
Qу
       1096 MWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKI 1155
           Db
        147 LWSLTYIASWFSGFTLAILGLLGVFSVPKVYESNQEAIDPHLATISGHLKNVQNIIDEKL 206
       1156 PGLK 1159
QУ
           | |:
        207 PFLR 210
Db
```

```
RESULT 9
A47283
calphotin - fruit fly (Drosophila melanogaster)
C; Species: Drosophila melanogaster
C;Date: 21-Sep-1993 #sequence revision 25-Apr-1997 #text change 21-Jul-2000
C; Accession: A47283
R; Ballinger, D.G.; Xue, N.; Harshman, K.D.
Proc. Natl. Acad. Sci. U.S.A. 90, 1536-1540, 1993
A; Title: A Drosophila photoreceptor cell-specific protein, calphotin, binds
calcium and contains a leucine zipper.
A; Reference number: A47283; MUID: 93165730; PMID: 8434015
A; Contents: photoreceptor cells
A; Accession: A47283
A; Status: preliminary
A; Molecule type: nucleic acid
A; Residues: 1-873 <BAL>
A; Cross-references: GB:L05080; NID:q157071; PIDN:AAA28420.1; PID:q157072
A; Note: sequence extracted from NCBI backbone (NCBIN: 124958, NCBIP: 124959)
C; Superfamily: collagen alpha 1(I) chain; fibrillar collagen carboxyl-terminal
homology; von Willebrand factor type C repeat homology
 Query Match
                       5.2%; Score 304.5; DB 2; Length 873;
 Best Local Similarity 21.9%; Pred. No. 3.1e-05;
 Matches 213; Conservative 126; Mismatches 362; Indels 273; Gaps
         62 AAGLSAAAVPPAAAAPLLDFSSDSVPPAPR--GPLPAAP----PAAP----ERQPSWER 110
Qу
            | | |
Db
         11 SAPVAAPVTPSAVAAPVQVVSPAAVAVAPAVVAPAPAAPTAVTPVAPPPTLASVOPATVT 70
         111 SPAAPAP----SLPPAAAVLPSKLPEDDEPPARPPPPPP------AGASPLAEPAAPP 158
Qу
                    Db
         71 VP-APAPIAAASVAPVASVAPPVVAAPTPPAASPVSTPPVAVAQIPVAVSAPVAPPVAAT 129
        159 STPAAPKRRGSGSVDETLFALP--AASEP----VIPSSAEKIMDLMEQPGNT---VSSGQ 209
Qу
             Db
        130 PTPVAPI-----PVAAPVIATPPVAASAPTPAAVTPVVSPVIATPPVVPANTTVPVAAPV 184
        210 EDFPSVLLETAASL-PSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERA 268
Qу
               185 AAVPAAVPVVAPVLAPAVAPAVAPVVAETPAPPPVAEIPVA--TIPECVAPLIPEVSVVA 242
Db
        269 TNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALH 328
Qу
                                              | [:|
        243 TKPLA-----AAEPVVVAPPATETPVVAPAAA 269
Db
        329 SPQESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDV 388
Qу
                      ::::
        270 SPHVS------ VAP----- 287
Db
Qу
        389 LAARANVESKVDRKCLEDSLEQKSL----GKDSEGRNEDASFPSTPEPVKDSSRAYITCA 444
               Db
        288 ---SASTEPPVAAATLTTAPETPALAPVVAESQVAANTVVATPPTPAPEPETIAPPVVAE 344
Qу
        445 SFTSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVODSEA 504
            : |:: | |:: |:|:
```

```
Db
        345 TPEVASVAVAETTPPVVPPVAAES-----IPAPVVATTPVPATLAVTDPD- 389
QУ
        505 DYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQ 564
                    Dh
         390 ------VTASAVPELPPVIAPSPVPSA------VAETPVDLA-----P 420
        565 ESLYP-TAQLCPSF--EEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAP-- 619
Qу
             421 PVLPPVAAEPVPAVVAEETPETPAPASAPVTI-AALD--IPEVAPVIAAPSDAPAEAPSA 477
Db
        620 -- PPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACD 677
Qу
                                                |:| || || :: :
        478 AAPIVS-----TPPT--
Db
                                            ----TASVPETTAPPAAVPTE 504
Qу
        678 ----LIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAE--LVEDSSPESEPVDLFSD-DS 729
                505 PIDVSVLSEAAIETPVAPPVEVTTEVAVADVAPPEAAEDLIIEPVEPPAPIPDLLEQTTS 564
Db
        730 IPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDAA 789
Qу
            Db
        565 VPAVEAAESTSSPIPETSLPPPNEAVA---SPEVAVAPITAPEPIPEP-EPSLATPTEPI 620
        790 SNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFV 849
Qу
             : |
                                   621 PVEAPV------VIQEAVDAVEVPVTETSTSIPETTVEFPEAV 657
Db
       850 SAKDDSPKLAKEYTDLEV-SDKSEIANIQSGAD----SLPCLELPCDLSFKNIYPKDEVH 904
QУ
            658 AEKVLDPAI----TEAPVTTQEPDVANINDGAPATEITTPAVEI------VT 699
Db
Qу
        905 VSDEFSENRSSVSKASISPSNVSALE-PQTEM--GSIVKSKSLTKEAEKKLPSDTEKEDR 961
            700 AAAEVSDTAIPLIDPPV-PQEIAVAEIPETETKPAEVIVEQS-TIPIEAPVPEVSKYAEP 757
Db
        962 SLSAVLSAELSKTS 975
Qу
            :| :||: |:
Db
        758 VISEAPAAEVPITA 771
RESULT 10
T24583
hypothetical protein T06D8.1 - Caenorhabditis elegans
C; Species: Caenorhabditis elegans
C;Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 15-Oct-1999
C; Accession: T24583
R; Palmer, S.
submitted to the EMBL Data Library, April 1995
A; Reference number: Z19909
A; Accession: T24583
A;Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-1829 <WIL>
A; Cross-references: EMBL:Z49130; PIDN:CAA88964.1; GSPDB:GN00020; CESP:T06D8.1
A; Experimental source: clone T06D8
C; Genetics:
A;Gene: CESP:T06D8.1
A; Map position: 2
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A;Introns: 1391/3; 1432/3; 1470/3; 1505/1; 1520/1; 1616/1; 1644/1; 1687/3; 1742/1

	Query Ma Best Loc Matches	al S	5.2%; Score 302.5; DB 2; Length 1829; Similarity 21.1%; Pred. No. 0.00011; 3; Conservative 157; Mismatches 469; Indels 245; Gaps	38
Qγ	7	2	EDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEE	42
Dh)	212	: :: :	271
Qу	7	43	DEEEDDEDLEELEVLERKPAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPP	99
Dk)	272		322
Qу	•	100	AAPERQPSWERSPAAPAPSLPPAAAVLPSKLPEDDE-PPARPPPP	143
Dk)	323	:	378
Qу	7	144	PPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMD	196
Dh)	379	: :	432
Qу	•	197	LMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEH : :: :: : : :	237
Db)	433	VTEASGEETTTAAATEASEETTTSAVTEGSGEDTTVVAVVESSGEQPASSSTSIPT	488
Qу	•	238	GYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPK : : : : : : :	297
Db)	489	-ELSKDDQVTEASGEETTTAAATEASEETTTSAVTEGSGEETTV-VAVVESSGEEPAS	544
Qу	-	298	GESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDI:: : : : : ::	350
Db)	545	SSTSIPTELSKDDKVTEASGEETTTAAATDASSEETTTSAVTEGSGEETTVVAVVESSD-	603
Qу		351	FNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQ	410
Db	1	604	: : : : : : : : :	655
Qу		411	KSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKT :: : : : : : : : : : : : :	470
Db		656	TTVVAVVESSGEEPASSSTSIPTELSKDDKVTEASGEETTTAAATDASSEETTTSAV	712
Qу		471	DEKKIEERKAQIITEKTSPK-TSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTP	529
Db			TEGSGEETTVVAVVESSDEEPASSSTSIPTELSKDDQVTEASGEETTTAAATEASEETTT	
Qу			DLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSF	
Db			SAVTEGSGEETTVVAVVESSGEEPASSSTSIPTELSKD-DQVTEASGEETTTAAATE	
Qу			EEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPY : :	637
Db				871
Ωу		638	EEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSE : : : : : : : : : : : : : : : :	697

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Db
         872 -----KDDQVTEASGEETTT---AAATEASEETTTSAVTEGSGEETTV 911
         698 IAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEV---PQTQEEAVMLMKESLTEVSET 754
QУ
             912 VAVVESSGEEPA----SSSTSIPTELSKDDQVTEASGEETTTAAATEASSEETTTSAVT 966
Db
         755 VAQHKEERLSASPQELGKPYLESFQP----- 791
Qу
                    : |::::|
         967 EGSGEETTTSAVTEGSGEETTTSAVPEGENSTTEAPAFVTGSEIEIPSSEESSSTTTHDP 1026
Db
         792 DIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESET-FSDSSPIEIIDEFPTFVS 850
Qу
              1027 SIPVITPKPSVSSTIENVMSKTSSEE---AAEKKIIGEHQTGKDDDAGKEDEDNMPAFVT 1083
Dh
Qу
                             --KDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCD 891
                               : | : | : | | |
                                            : : : : [
Db
        1084 ANPAGTSTTESAENVTSTGEEDENIKMAKELGKQFAADLAKLA----- 1126
Qу
         892 LSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKK 951
                    1127 -----AKDGVNLT-ETADAKDSGETAHVEDEQVSSTE--SSIGSEETTTTVNKETTEE 1176
Db
         952 LPSDTEKEDRSLSAVLSAELSKTS 975
Qу
                : : | : : | | :
        1177 HHEASGEEDDAPAFVTGAPTDSTT 1200
Db
RESULT 11
calcium-binding protein calphotin - fruit fly (Drosophila melanogaster)
C; Species: Drosophila melanogaster
C;Date: 21-Sep-1993 #sequence revision 18-Nov-1994 #text change 21-Jul-2000
C; Accession: A47282
R; Martin, J.H.; Benzer, S.; Rudnicka, M.; Miller, C.A.
Proc. Natl. Acad. Sci. U.S.A. 90, 1531-1535, 1993
A; Title: Calphotin: a Drosophila photoreceptor cell calcium-binding protein.
A; Reference number: A47282; MUID: 93165729; PMID: 8094559
A; Accession: A47282
A; Status: preliminary
A; Molecule type: nucleic acid
A; Residues: 1-865 < MAR>
A; Cross-references: GB:L02111; NID:g157031; PIDN:AAA28405.1; PID:g157032
A; Experimental source: photoreceptor cells
A; Note: sequence extracted from NCBI backbone (NCBIN:124955, NCBIP:124956)
C; Genetics:
A; Gene: FlyBase: Cpn
A; Cross-references: FlyBase: FBgn0010218
C; Superfamily: collagen alpha 1(I) chain; fibrillar collagen carboxyl-terminal
homology; von Willebrand factor type C repeat homology
C; Keywords: calcium binding
 Query Match 5.1%; Score 295.5; DB 2; Length 865; Best Local Similarity 21.3%; Pred. No. 7.1e-05;
 Query Match
 Matches 217; Conservative 127; Mismatches 379; Indels 295; Gaps
Qу
         62 AAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPER----QPSWERSPAAPAP 117
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Db	11	SAPVAAPVTPSAVAAPVQVVSPAAVAPAPAAPIAVTPVAPPPTLASVQPATVTIP-APAP	69
Qy	118	SLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTP	161
Db	70	: : :	129
Qy	162	-AAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQED	211
Db	130	: :: VAAPVIATPPVAASAPTPAAVTPVISPVIASPPVVPANTTVPVAAPVAA	178
Qу	212	FPSVLLETAASL-PSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATN : : :	270
Db	179	VPAAVPVVAPVLAPAVAPAVAPVVAETPAPPPVAEIPVATIPECVAPLIPEVSVVATK	236
Qy	271	PFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSP	330
Db	237		263
Qy	331	QESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLA	390
Db	264	HVSAVETAVVAPV	279
Qy	391	ARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASF :	446
Db	280	-SASTEPPVAAATLTTAPETPALAPVVAESQVAANTVVATPPTPAPEPETIAPPVVAETP	338
Qy	447	TSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADY : : :: : : : :	506
Db	339	EVASVAVAETTPPVVPPVAAESIPAPVVATTPVPATLAVTDPD	381
Qу	507	VTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQES	566
Db	382	VTASAVPELPPVIAPSPVPSAVAETPVDLAPPV	414
Qу	567	LYP-TAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAP	619
Db	415	LPPVAAEPVPAVVAEETPETPAPASAPVTI-AALDIPEVAPVIAAPSDAPAEAPSAAA	471
Qy	620	PPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACD	677
Db	472	PIVSTPPTTASVPETTAPPAAVPTEPI	498
Qy	678	LIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSD-DSIP :: : : : : : : :	731
Db	499	DVSVLSEAAIETPVAPPVEVTTEVAVADVAPPEAAADLIIEPVEPPAPIPDLLEQTTSVP	558
Qy	732	EVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDAASN : : : : : :	791
Db	559	AVEAAESTSSPIPETSLPPPNEAVASPEVAVAPITAPEPIPEP-EPSLATPTEPIPV	614
Qy		DIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSA: :: :: : : :	
Db	615	EAPVVIQEAVDAVEVPVTETSTSIPETTVEFPEAVAE	651
Qу	852	KDDSPKLAKEYTDLEV-SDKSEIANIQSGADSLPCLELPCDLSFKNIYP-	899
Db	652	KVLDPAI TEAPVTTOEPDVANINDGAPATEITTPAVEIVTAAAEVSDIAIPVIDPP	707

```
900 - KDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMG------ 936
Qу
               Db
        708 VPQEIAVA-EIPETDTKPAEVIVEQSTIPIEAPVPEVSKYAEPVISEAPAAEVPITAGDN 766
        937 ----SIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGV 990
ОУ
               767 PDNTSVGISEVVPTIAEKPVEEVPTSEIPEQSSSPSDSVPVAKITPLL--RDLOTTDV 822
RESULT 12
T34513
hypothetical protein ZK783.1 - Caenorhabditis elegans
C; Species: Caenorhabditis elegans
C; Date: 29-Oct-1999 #sequence revision 29-Oct-1999 #text change 29-Oct-1999
C; Accession: T34513
R; Favello, A.; Vaudin, M.
submitted to the EMBL Data Library, August 1994
A; Description: The sequence of C. elegans cosmid ZK783.
A; Reference number: Z21536
A; Accession: T34513
A; Status: preliminary; translated from GB/EMBL/DDBJ
A; Molecule type: DNA
A; Residues: 1-3507 <FAV>
A;Cross-references: EMBL:U13646; PIDN:AAC24418.1; GSPDB:GN00021; CESP:ZK783.1
A; Experimental source: strain Bristol N2; clone ZK783
C; Genetics:
A; Gene: CESP: ZK783.1
A; Map position: 3
A;Introns: 14/1; 48/2; 84/1; 196/3; 303/1; 381/1; 586/1; 605/1; 1175/3; 1207/1;
1409/2; 1450/1; 1760/1; 1898/1; 2153/1; 2628/1; 2681/1; 2899/1; 2946/1; 3088/3;
3184/1; 3283/1; 3346/3; 3365/3; 3484/3; 3504/1
 Query Match
                       5.0%; Score 292; DB 2; Length 3507;
 Best Local Similarity 20.6%; Pred. No. 0.00071;
 Matches 226; Conservative 146; Mismatches 491; Indels 236; Gaps
Qу
         11 SSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEEDEEDDEDL--EELEVLERKPAAGLSAA 68
            2043 SSSSEAPLTSSPATTTEVITESSVKSTTPKEESSSEITVKLSSKSPEVTESSVKSSPSTP 2102
Db
         69 AVPPAAAAPLLDFSSDSVPPAPRGPLPAAPP----AAPERQPSWERSPA----APAPSL 119
Qу
           Db
       2103 STTSQSVTSTVPETSKSTVLSSEAPVTSTSPTEVHTSSETKPSLSASSTTGDTNSTTPST 2162
        120 PPAAAVLPSKLPEDDEPPARPPPPPPAGASP-LAEP-----AAPPSTPAAPKRRGSGSV 172
QУ
              Db
       2163 SSLASVKSTSAPEGTS--ASVAPVKLSSLSPDVSQPSTKTFDATESSTVQASETSSGTSV 2220
QУ
        173 DET-----LFALPAASEPVI-PSSAEKIMDLMEOPGNTVSSGOEDFP----SV 215
                          2221 KSTSEPESHVTKLSITSSNPSSSVPVTSPKSTPTVPESTEQPTSTTPSGQSLTPMNSNSE 2280
Db
Qу
        216 LLETA---ASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERAT--N 270
                      Db
       2281 VLTTSEPHVLSSSLSPDVSQSSTTPNNLSESSTVETPKTSSEVSLNSEEPSTTEAPTTLS 2340
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Qy		PFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKD	317
Db .		PDILSTTTNNLSQSSTVSTEDRSEISSENSEKPTSAPELVTSSVTHVASSSPDVPTESSE	2400
Qу	318	KEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFK	370
Db	2401	: : : : : : : : :	2460
Qy.	371	PFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTP	430
Db	2461	: : : : : ESSSKSPVSSSTEGISVVTSTEFSKVPESTISSVLEEDLT-KTTP	2504
Qy	431	EPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEE	477
Db	2505	: : : : : : : : : : : : SPILEETTTASETSEPLTEDSLTVSVRIHELTTSSENVPKESESTTTSSESSKPSQ	2560
Qу	478	RKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLV	532
Db	2561	:	2618
Qу	533	QEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPD	590
Db	2619	:: : :: : TSSEPSESTKRTTVSTTVSTTTPTEETTTSESLILTAAPSKPTESTTESSEAPTTP-	2674
Qу		IVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGT	
Db	2675	:	2692
Qу	651	KEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAE	710
Db	2693	::	2738
Qу	711	LVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKE-ERLSASPQE	769
Db	2739	: : : ::: : : : NSSTSSPTSSEASVKLSSLFPESITSEAVTVSSRAPAEITMSSESHREISTVSSEPSE	2796
Qу	770	LGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSS	822
Db	2797	: : : : : EIPLSTTVSPNVVTASSIPSEE-PILSSVTSSSTPRVRLITGTPDDLIVSVTVPSHG	2853
Qу	823	-KEDKIKESETFSDS-SPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGA : : : :	880
Db	2854	NRRQNITASSVPSNSTSPIILPSESLTTPQPPPTTTTTAKPATTSGK	2900
Qу	881	DSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVK	940
Db	2901	:: :: ::	2929
Qy	941	SKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLL	1000
Db	2930	: ::: : :: : : : : :: : : : : : : : :	2976
Qу	1001	SLTVFSIVSVTAYIALALL 1019	
Db	2977		

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RESULT 13
A56577
microtubule-associated protein MAP 1B - rat (fragment)
C; Species: Rattus norvegicus (Norway rat)
C;Date: 21-Jul-1995 #sequence_revision 21-Jul-1995 #text change 16-Feb-1997
C; Accession: A56577
R; Zauner, W.; Kratz, J.; Staunton, J.; Feick, P.; Wiche, G.
Eur. J. Cell Biol. 57, 66-74, 1992
A; Title: Identification of two distinct microtubule binding domains on
recombinant rat MAP 1B.
A; Reference number: A56577; MUID: 92347374; PMID: 1639092
A; Accession: A56577
A; Status: preliminary
A; Molecule type: mRNA
A; Residues: 1-2364 <ZAU>
A; Cross-references: GB:X60550
A; Experimental source: brain
A; Note: nucleotide sequence not given; conceptual translation not complete
C; Superfamily: microtubule-associated protein MAP1B
 Query Match
                      5.0%; Score 291.5; DB 2; Length 2364;
 Best Local Similarity 20.0%; Pred. No. 0.00043;
 Matches 220; Conservative 167; Mismatches 456; Indels 255; Gaps 44;
         30 TEPEDEEDEEEEDDEDLEELEVLERKPAAGLSAAAVPPAAAAPLLDFSSDSVPPA 89
Oy
           Db
        913 SEEEGEEEEDKAEDAREEDHEPDKTE----AEDYVMAVVDKAAEAGVTEDQYDFL--- 963
         90 PRGPLPAAPPAAPERQPSWERSPAAPAPSLPPAAAVLPSKLPEDDEPPARPPPPPPAGAS 149
QУ
                    964 -----AS 999
Db
Qу
        150 PLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMDLMEOPGN---TVS 206
                 1000 DEENREDQPEEFTAT----SGYTQST---IEISSEPTPMDEMSTPRDVMTDETNNEETES 1052
Db
Qу
        207 SGQE-----DFPSVLLETAASLP---SLSPLS----TVSFKEHGYLGNLSAVSSSEGTIE 254
             Db
       1053 PSQEFVNITKYESSLYSQEYSKPVVASFNGLSDGSKTDATDGRDYNASASTISPPSSMEE 1112
Qу
        255 ETLNEAS------KELPERATNPFVNRDLAEF--SELEYSEMG--- 289
                                 Db
       1113 DKFSKSALRDAYRPEETDVKTGAELDIKDVSDERLSPAKSPSLSPSPPSPIEKTPLGERS 1172
        290 -----SSFKGSPKGESAILVENTKEEVIVR----SKDKEDL------VCSAALHSP 330
Qу
                 Db
       1173 VNFSLTPNEIKASAEGEATAVVSPGVTQAVVEEHCASPEEKTLEVVSPSQSVTGSAGHTP 1232
        331 -QESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFK-----PFEQAWEVKD 380
Qу
             1233 YYQSPTDEKSSHLPTEVT-----ENAQAVPVSFEFTEAKDENERSSISPMDE--PVPD 1283
Db
        381 TYEGSRDVLAARANVESKVDRKCLED--SLEQKSLGKDS----EGRNEDASFPSTPEPVK 434
QУ
           : ||: :
                               Db
       1284 SESPIEKVLSPLRSPPLIGSESAYEDFLSADDKALGRRSESPFEGKNGKQGFSDKESPVS 1343
        435 DSSRAYITCASFTSATESTTANTFPLLEDHTSENKT------DEKKI---- 475
QУ
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Db	1344	: : : : : : DLTSDLYQDKQEEKRAGFIPIKEDFSPEKKASDAEIMSSQSALALDERKLGGDGS	1398				
Qy	476	EERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAV	520				
Db	1399	:	1455				
Qy	521	SNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQES-LYPTAQLCPS	576				
Db	1456	: : : : :: :	1514				
Qу	577	FEEAEATP	610				
Db	1515	:: :: : : : : : : : : : : :	1574				
Qу	611	PSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFN	661				
Db	1575	:: : ::: : : :	1630				
Qy	662	AAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEP	721				
Db	1631	:: : : : : : :	1680				
Qу	722	VDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYL-ESFQP	780				
Db	1681	: :: :: : : :: : : SPLTPRESSPTYSPGFSDSTSGAKES-TAAYQTSSSPPIDAAAAEPYGFRSSMLFDTMQH	1739				
Qу	781	NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	840				
Db	1740	: :: :: : : :: : : :: HLALSRDLTTSSVEKDNGGKTPGDFNYAYQKPESTTESPDEEDYDYESHEKTIQ	1793				
Qу	841	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKN : : :	896				
Db	1794	: :: : : :: AHDVGGYYYEKTERTIKSPCDSGYSYETIEKTTKTPEDGGYSCEITEKT	1842				
Qу	897	IYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSL	944				
Db	1843	: : : : : : : : : TRTPEEGGYSYEISEKTTRTPEVSGYTYEKTERSRRLLDDISNGYDDTEDGGHTLGDCSY	1902				
Qу	945	TKEAEKKLPSDTEKEDRS 962					
Db	1903	: : : SYETTEKITSFPESESYS 1920					
RESULT 14 T19431 hypothetical protein C25A1.10 - Caenorhabditis elegans C;Species: Caenorhabditis elegans C;Date: 15-Oct-1999 #sequence_revision 15-Oct-1999 #text_change 21-Jan-2000 C;Accession: T19431 R;Mortimore, B. submitted to the EMBL Data Library, October 1996 A;Reference number: Z19124 A;Accession: T19431 A;Status: preliminary; translated from GB/EMBL/DDBJ A;Molecule type: DNA A;Residues: 1-971 <wil></wil>							

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A;Cross-references: EMBL:Z81038; PIDN:CAB02755.1; GSPDB:GN00019; CESP:C25A1.10
A; Experimental source: clone C25A1
C;Genetics:
A; Gene: CESP: C25A1.10
A; Map position: 1
A; Introns: 38/3; 92/3; 201/3; 919/3
C; Superfamily: nucleolus-cytoplasm shuttle phosphoprotein
  Query Match
                     4.9%; Score 288.5; DB 2; Length 971;
 Best Local Similarity 20.6%; Pred. No. 0.00016;
 Matches 198; Conservative 127; Mismatches 391; Indels 247; Gaps
                                                              35:
          7 SSLVSSSTDSPPRPPPAFKYQFVTEP---EDEEDEEEEEDEEEDDEDLEELEVLERKPAA 63
Qу
           147 SSSDSDSDDEPPKKAPAVTTKVAPKPMAKKQDTSDSDSDSDSDSDSKSKKANPVKVTPVA 206
Db
         64 GLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAP-----EROPSWE 109
Qу
               Db
        207 NVLQKVVAKKAASSSSDSSDDEKKPAAK-PTPAKPTPKPVVKKAESSSDSSDDEKKPVAK 265
        110 RSPAAPAPSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGS 169
Qу
            Dh
        266 PAPAKATPK-PAAKKADSSSDSSDDEAPAK-KTPAKAAPKPVAKKAESSSDSSDDEKK-- 321
        170 GSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPL 229
Qу
                   Db
        322 -----PAAK----PTPAKATPKPVAKKAESSSDSSDDEKKPVAKPAPAKATPKPV 367
        230 STVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSE-- 287
Qу
                       368 AK------KAESSSDSSDDEK-KPAAKPTPAKATPKPVAKKAESSSDSSDDEKK 414
Db
        288 ---MGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSP 344
Qу
               : | | : | |
                                  415 PVAKPTSAKATPK-----PAAKKAD---SSSDSSDDEAPAKKTPAKAAP 455
Db
        345 E---KTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRD------VLAARANV 395.
Qу
           : | : :: : : | | | : | : | |
                                                   . >
        456 KPASKKAESSSDSSDDEKPAAKSTPAKITPKPTAKKVASSSSDSSDDEKKPAAKPTPANA 515
Db
        396 ESKVDRKCLEDSL----EQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSAT 450
Qу
                         516 TPKPVAKKAESSSDSSDDEKKPVAKPTSAK-----ATPKPAAKKADLSSDFSDDEAPA 568
Db
        451 ESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPK-TSNPFLVAVQDSEADYVTT 509
QУ
                              |: |:
                       : | : : |
                                      Db
        569 KKTPAKAAPKPASKKAESSSDSSDDEKPAAKSTPAKTTPKPTAKKAASSSSDSSDD--EK 626
        510 DTLSKVTEAAVSNMPEGLTPDLVQEACESELN-----EATGTKIAYETKVDLVQTSEAI 563
Qу
             627 KPVAKPTSAKATPKPAAKKADSSSDSSDDEAPAKKTPVKPTPVKIVAK-KVD--SSSDSS 683
Db
        564 QESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVOPSVSPLEAPPPVS 623
Qу
                        684 DDEKKPT------SATPVKVTP-----S 710
Db
        624 YDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETK 683
Qу
```

```
1:
                                                          : :
Db
          711 SDS--SDDEKKPVVKQTPNVV------PKKEKAASSSDDSS-----DDEKK 748
          684 LSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVML 743
Qу
               : : | : |
                                 |: |: :
                                            | \cdot |
                                                        111
Db
          749 PTAKPTP-
                              ---KATPKQSAKKADSSDDS-----SDDEAPA----- 779
Ov
          744 MKESLTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKIS 803
Db
                                                      -KKTPAKSTPAKTAVKKEASS 799
          804 LQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS--PKLAKE 861
Qу
                         |\cdot|
                                 :: | : | : | : | : | : | : | : |
Db
                        -SSDDSSDDEKTKKKSATTPAKSTPKTALKKAESSDSSDDDEDLPKPSKA 848
Qу
          862 YTDL-EVSDKSEIANIQSGADSLPCLEL-PCDLSFKNIYPKDEVHVSDEFSENRSSVSKA 919
                  Db
          849 VTPRPQRADSEESAETEESSSRTPALKAKPLATSTEKAVYENRKRKSSPF--RRVQMTKD 906
          920 SIS 922
Qу
              1:1
Db
          907 SVS 909
RESULT 15
ORMSP1
microtubule-associated protein MAP1B - mouse
N; Alternate names: microtubule-associated protein MAP1(X); microtubule-
associated protein MAP1.2; microtubule-associated protein MAP5
C; Species: Mus musculus (house mouse)
C;Date: 30-Sep-1991 #sequence revision 30-Sep-1991 #text change 01-Sep-2000
C; Accession: S07549; S44387; A33645
R; Noble, M.; Lewis, S.A.; Cowan, N.J.
J. Cell Biol. 109, 3367-3376, 1989
A; Title: The microtubule binding domain of microtubule-associated protein MAP1B
contains a repeated sequence motif unrelated to that of MAP2 and tau.
A; Reference number: A33645; MUID: 90094539; PMID: 2480963
A; Accession: S07549
A; Molecule type: mRNA
A; Residues: 1-2464 < NOB>
A;Cross-references: EMBL:X51396; NID:g52999; PIDN:CAA35761.1; PID:g53000
R; Sanchez, C.; Padilla, R.; Paciucci, R.; Zabala, J.C.; Avila, J.
Arch. Biochem. Biophys. 310, 428-432, 1994
A; Title: Binding of heat-shock protein 70 (hsp70) to tubulin.
A; Reference number: S44387; MUID: 94234720; PMID: 8179328
A; Accession: S44387
A; Status: preliminary
A; Molecule type: protein
A; Residues: 653-663, 'IC' <SAN>
C; Superfamily: microtubule-associated protein MAP1B
C; Keywords: microtubule binding; phosphoprotein; tandem repeat
F;589-786/Domain: microtubule binding #status experimental <MTB>
F;589-592,639-642,649-652,655-658,660-663,668-671,674-677,679-682,683-686,687-
690,691-694,695-698,699-702,708-711,712-715,716-719,720-723,727-730,758-761,764-
767,783-786/Region: 4-residue repeats (K/R-K-E/D-X)
F;1861-2064/Region: 17-residue repeats
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F; 91, 116, 351, 888, 1124, 1153, 1168, 1208, 1662, 1877, 1918, 2003, 2030, 2054, 2083/Binding
site: phosphate (Ser) (covalent) #status predicted
F;147,969,1336,1562,1563,1702,1708,1990,2057,2063,2419/Binding site: phosphate
(Thr) (covalent) #status predicted
F;1953/Binding site: phosphate (Tyr) (covalent) #status predicted
 Ouery Match
                     4.9%; Score 284; DB 1; Length 2464;
 Best Local Similarity 20.7%; Pred. No. 0.00092;
 Matches 233; Conservative 159; Mismatches 424; Indels 312; Gaps
                                                            51:
Qу
         31 EPEDEEDEEEEDEEED-----DEDLEELE-----VLERKPAAG-----LSAAAVP 71
           1009 EAEQSEEGEEDKAEDAREEGYEPDKTEAEDYVMAVADKAAEAGVTEEQYGYLGTSAKO 1068
Db
         72 PAAAAPLLDFSS----DSVPPAPRGPLPAAPPAAPERQP-----SWERSPAAPA 116
Qу
              Db
       1069 PGIQSPSREPASSIHDETLPGGSESEATASDEENREDQPEEFTATSGYTQSTIEISSEPT 1128
        117 P---SLPPAAAVLPSKLPEDDEPPARP------PPPPPAGASPLAEPA-- 155
Qу
           1129 PMDEMSTPRDVMSDETNNEETESPSQEFVNITKYESSLYSQEYSKPAVASFNGLSEGSKT 1188
Db
        156 -----APPSTPAAPKRRGSGSVDETLFALPA-----ASEPVIPSSAEKIMDLMEQ 200
Qу
                  1189 DATDGKDYNASASTISPP----SSMEEDKFSKSALRDAYCSEEKELKASAE--LDIKDV 1241
Db
        201 PGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEA 260
Qу
              | | | | |
       1242 SDERLS-----PAKSPSLSP------SPPSPIEKT---- 1265
Db
        261 SKELPERATNPFVNRDLAEFS----ELEYSEMGSSFKGSPKGESAILVEN--TKEEVIVR 314
Qу
             1266 -- PLGERSVN-----FSLTPNEIKVSAEGEARSVSPGVTQAVVEEHCASPEEKTLE 1314
Db
        315 SKDKEDLVCSAALHSP-QESPVGKEDRVVSPEKTMDIFNEMOMSVVA-PVREEYADFK-- 370
Qу
                Db
       1315 VVSPSQSVTGSAGHTPYYQSP-----TDEKSSHLPTEVSENAQAVPVSFEFSEAKDE 1366
        371 -----PFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLED--SLEQKSLGKDS---- 417
Qу
                † ::   | |:   | |: | ||: |
Db
       1367 NERASLSPMDE--PVPDSESPVEKVLSPLRSPPLLGSESPYEDFLSADSKVLGRRSESPF 1424
        418 EGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKT----- 470
Qу
                Db
       1425 EGKNGKQGFPDRESPVSDLT----STGLYQDKQEEKSTGFIPIKEDFGPEKKTSDVETMS 1480
        471 ------DEKKI-------EERKAQIITEKTSPKTSNPFLVAVQDSEAD 505
Qу
                                  Db
       1481 SQSALALDERKLGGDVSPTQIDVSQFGSFKEDTKMSISEGTVSDKSATPVDEGV--AEDT 1538
        506 YVTTDTLSKVTEAAV--SNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAI 563
QУ
           Db
       1539 YSHMEGVASVSTASVATSSFPEPTTDD-VSPSLHAEVGSPHSTEVDDSLSVSVVOTPTTF 1597
       564 QES-LYPTAQLCP---SFEEAEATP-------SPVLPDIVMEAPLNSLL-- 601
Qу
       1598 QETEMSPSKEECPRPMSISPPDFSPKTAKSRTPVQDHRSEQSSMSIEFGQESPEHSFAMD 1657
Db
```

Qу	602	PSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVA	644
Db	1658	FSRQSPDHPTLGASVLHITENGPTEVDYSPCDIQDSSLSHKIPPTEEPSYTQDNDLS	1714
Qy	645	LKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKS : : : : : :	704
Db	1715	-ELISVSQVEASPSTSSAHTPSQIASPLQEDTLSDVVPPREMSLYASLA	1762
Qу	, 705	VPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKE : : :	760
Db	1763	SEKVQSLEGEKLSPKSDISPLTPRESSPLYSPGFSDSTSAAKETAAAH	1810
Qу	761	ERLSASPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFN : : : : : : : :	810
Db		-QASSSPPIDAATAEPYGFRSSMLFDTMQHHLALNRDLTTSSVEKDSGGKTPGDFN	
Qу		TAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLE :: ::: : :: : :	
Db		YAYQKPENAAGSPDEEDYDYESQEKTIRTHDVVRYYYEKTERTIKSPCDSGYSYETIE	
Qу		VSDKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKAS : : : : :	
Db		KTTKTPEDGGYTCEITEKTTRTPEEGGYSYEISEKTTRTPEVSGYTYEK	1972
Qу		ISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRS 962 : : : : : :	
Db	1973	TERSRRLLDDISNGYDDTEDGGHTLGDCSYSYETTEKITSFPESESYS 2020	

Search completed: December 19, 2003, 15:37:23 Job time : 34 secs

> GenCore version 5.1.6 Copyright (c) 1993 - 2003 Compugen Ltd.

OM protein - protein search, using sw model

Run on: December 19, 2003, 15:29:47; Search time 18 Seconds

(without alignments)

3038.448 Million cell updates/sec

Title: US-09-830-972-2

Perfect score: 5848

Sequence: 1 MEDIDQSSLVSSSTDSPPRP.....VKDAMAKIQAKIPGLKRKAD 1163

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 127863 seqs, 47026705 residues

Total number of hits satisfying chosen parameters: 127863

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

SwissProt 41:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result		% Query				
No.	Score	Match	Length	DB	ID	Description
, 1	5848	100.0	1163	1	RTN4_RAT	Q9jk11 rattus norv
2	4403.5	75.3	1192	1	RTN4_HUMAN	Q9nqc3 homo sapien
3	917	15.7	199	1	RTN4_MOUSE	Q99p72 mus musculu
4	801	13.7	777	1	RTN1_RAT	Q64548 rattus norv
5	789.5	13.5	776	1	RTN1_HUMAN	Q16799 homo sapien
6	625.5	10.7	236	1	RTN3_HUMAN	O95197 homo sapien
7	625.5	10.7	237	1	RTN3_MOUSE	Q9es97 mus musculu
8	509	8.7	471	1	RTN2_MOUSE	070622 mus musculu
9	490	8.4	545	1	RTN2_HUMAN	075298 homo sapien
10	308.5	5.3	5147	1	PCLO_HUMAN	Q9y6v0 homo sapien
11	300	5.1	5120	1	PCLO CHICK	Q9pu36 gallus gall
12	295.5	5.1	865	1	CPN_DROME	Q02910 drosophila
13	291.5	5.0	2459	1	MAPB_RAT	P15205 rattus norv
14	288	4.9	2468	1	MAPB HUMAN	P46821 homo sapien
15	284	4.9	2464	1	MAPB MOUSE	P14873 mus musculu
16	282.5	4.8	2805	1	MAPA HUMAN	P78559 homo sapien
17	279	4.8	5038	1	PCLO MOUSE	Q9qyx7 mus musculu
18	275.5	4.7	3924	1	ANK2 HUMAN	Q01484 homo sapien
19	275	4.7	1972	. 1	P531 HUMAN	Q12888 homo sapien
20	272	4.7	5085	1	PCLO RAT	Q9jks6 rattus norv
21	267	4.6	1781	1	AK12_HUMAN	Q02952 homo sapien
22	265.5	4.5	1828	1	MAP2 MOUSE	P20357 mus musculu
23	261.5	4.5	1087	1	NFH MOUSE	P19246 mus musculu
24	259.5	4.4	1020	1	NFH HUMAN	P12036 homo sapien
25	259.5	4.4	2404	1	SON MOUSE	Q9qx47 mus musculu
26	258.5	4.4	1861	1	MAP2 RAT	P15146 rattus norv
27	257.5	4.4	2715	1	TRX2_HUMAN	Q9umn6 homo sapien
28	256	4.4	1723	1	AIM1 HUMAN	Q9y4k1 homo sapien
29	255.5	4.4	1140	1	YM96 YEAST	Q04893 saccharomyc
30	254.5	4.4	6632	1	UN89 CAEEL	001761 caenorhabdi
. 31	254	4.3	1616	1	P200 MYCGE	Q49429 mycoplasma
32	254	4.3	3381	1	PGCV BOVIN	P81282 bos taurus
33	253	4.3	3421	1	TEGU HSVEB	P28955 equine herp
34	252.5	4.3	1411	1	TCOF HUMAN	Q13428 homo sapien
35	252	4.3	2738	1	PGCV RAT	Q9erb4 rattus norv
36	250.5	4.3	4377	1	ANK3 HUMAN	Q12955 homo sapien
37	248.5	4.2	1189	1	YJH6_YEAST	P47035 saccharomyc
38	248	4.2	1827	1	MAP2 HUMAN	P11137 homo sapien
3,9	245.5	4.2	1001	1	IF2 SYNY3	P72689 synechocyst
40	244	4.2	1338	1	ACIN_MOUSE	Q9jix8 mus musculu
41	241.5	4.1	8545	1	ANC1 CAEEL	Q9n4m4 caenorhabdi
42	238.5	4.1	831	1	NFH RAT	P16884 rattus norv
					_	= = 3001

```
      43
      238.5
      4.1
      1341
      1
      ACIN_HUMAN
      Q9ukv3 homo sapien

      44
      238
      4.1
      1306
      1
      MSB2_YEAST
      P32334 saccharomyc

      45
      236.5
      4.0
      2774
      1
      MAPA_RAT
      P34926 rattus norv
```

ALIGNMENTS

```
RESULT 1
RTN4 RAT
ID
     RTN4 RAT
                     STANDARD;
                                    PRT; 1163 AA.
     Q9JK11; Q9JK10; Q9R0D9; Q9WUE9; Q9WUF0;
AC
DT
     28-FEB-2003 (Rel. 41, Created)
DT
     28-FEB-2003 (Rel. 41, Last sequence update)
DT
     15-SEP-2003 (Rel. 42, Last annotation update)
DE Reticulon 4 (Neurite outgrowth inhibitor) (Nogo protein) (Foocen)
DE
     (Glut4 vesicle 20 kDa protein).
GN
     RTN4 OR NOGO.
OS
     Rattus norvegicus (Rat).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
     Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OC
OX
     NCBI TaxID=10116;
RN
     [1]
RΡ
     SEQUENCE FROM N.A. (ISOFORM 3), AND PARTIAL SEQUENCE.
RC
     STRAIN=Sprague-Dawley; TISSUE=Adipocyte;
RX
     MEDLINE=99249816; PubMed=10231557;
     Morris N.J., Ross S.A., Neveu J.M., Lane W.S., Lienhard G.E.;
RA
     "Cloning and characterization of a 22 kDa protein from rat adipocytes:
RT
     a new member of the reticulon family.";
RT
RL_{-}
     Biochim. Biophys. Acta 1450:68-76(1999).
RN
     SEQUENCE FROM N.A. (ISOFORMS 1; 2 AND 3).
RP
RX
     MEDLINE=20129258; PubMed=10667796;
RA
     Chen M.S., Huber A.B., Van der Haar M.E., Frank M., Schnell L.,
     Spillmann A.A., Christ F., Schwab M.E.;
RA
RT
     "Nogo-A is a myelin-associated neurite outgrowth inhibitor and an
RT
     antigen for monoclonal antibody IN-1.";
RL
     Nature 403:434-439(2000).
RN
     [3]
RP
     SEQUENCE FROM N.A. (ISOFORMS 2 AND 4).
RC
     STRAIN=Wistar Kyoto; TISSUE=Vascular smooth muscle;
RA
     Ito T., Schwartz S.M.;
     "Cloning of a member of the reticulon gene family in rat: one of two
RT
RT
     minor splice variants.";
     Submitted (FEB-1999) to the EMBL/GenBank/DDBJ databases.
RL
RN
     [4]
     FUNCTION.
RΡ
RX
     MEDLINE=22033691; PubMed=12037567;
RA
     GrandPre T., Li S., Strittmatter S.M.;
     "Nogo-66 receptor antagonist peptide promotes axonal regeneration.";
RT
RL
     Nature 417:547-551(2002).
     -!- FUNCTION: Potent neurite outgrowth inhibitor which may also help
CC
CC
         block the regeneration of the nervous central system in adults (By
CC
         similarity).
CC
     -!- SUBUNIT: Binds to RTN4R. Interacts with Bcl-xl and Bcl-2 (By
CC
         similarity).
```

-!- SUBCELLULAR LOCATION: Integral membrane protein. Anchored to the

CC

```
CC
         membrane of the endoplasmic reticulum through 2 putative
CC
         transmembrane domains (By similarity).
CC
     -! - ALTERNATIVE PRODUCTS:
CC
         Event=Alternative splicing; Named isoforms=4;
CC
         Name=1; Synonyms=Noqo-A, NI-220-250;
           IsoId=Q9JK11-1; Sequence=Displayed;
CC
CC
         Name=2; Synonyms=Nogo-B, Foocen-M1;
CC
           IsoId=Q9JK11-2; Sequence=VSP 005658;
CC
         Name=3; Synonyms=Nogo-C, VP20;
CC
           IsoId=Q9JK11-3; Sequence=VSP 005656, VSP 005657;
CC
         Name=4; Synonyms=Foocen-M2;
CC
           IsoId=Q9JK11-4; Sequence=VSP 005659;
CC
     -!- TISSUE SPECIFICITY: Isoforms 1, 2 and 3 are present in optic
CC
         nerve, spinal cord and cerebral cortex. Isoforms 1 and 2 are
CC
         present in dorsal root ganglion, sciatic nerve and PC12 cells
CC
         after longer exposure. Isoforms 2 and 3 are detected in kidney,
CC
         cartilage, skin, lung and spleen. Isoform 3 is expressed at high
CC
         level in skeletal muscle. In adult animals isoform 1 is expressed
CC
         mainly in the nervous system.
CC
     -!- SIMILARITY: Contains 1 reticulon domain.
CC
CC
     This SWISS-PROT entry is copyright. It is produced through a collaboration
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
     the European Bioinformatics Institute. There are no restrictions on its
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     use by non-profit institutions as long as its content is in no way
     modified and this statement is not removed. Usage by and for commercial
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     entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
     or send an email to license@isb-sib.ch).
CC
     ______
CC
DR
     EMBL; AF051335; AAF01564.1; -.
     EMBL; AJ242961; CAB71027.1; -.
DR
DR
     EMBL; AJ242962; CAB71028.1; -.
DR
     EMBL; AJ242963; CAB71029.1; -.
DR
     EMBL; AF132045; AAD31019.1; -.
DR
     EMBL; AF132046; AAD31020.1; -.
DR
     GO; GO:0030176; C:endoplasmic reticulum membrane, intrinsic p. . .; IDA.
     GO; GO:0005635; C:nuclear membrane; ISS.
DR
     GO; GO:0005515; F:protein binding activity; ISS.
DR
     GO; GO:0019987; P:negative regulation of anti-apoptosis; ISS.
DR
     GO; GO:0030517; P:negative regulation of axon extension; ISS.
DR
     InterPro; IPR003388; Reticulon.
DR
     Pfam; PF02453; Reticulon; 1.
     PROSITE; PS50845; RETICULON; 1.
DR
     Endoplasmic reticulum; Alternative splicing; Transmembrane.
KW
FT
     DOMAIN
                   1
                        989
                                 CYTOPLASMIC (Potential).
FT
     TRANSMEM
                 990
                       1010
                                 POTENTIAL.
FT
     DOMAIN
                1011
                      1104
                                 LUMENAL (Potential).
FT
    TRANSMEM
               1105
                      1125
                                 POTENTIAL.
FT
    DOMAIN
               1126
                      1163
                                 CYTOPLASMIC (Potential).
FT
    DOMAIN
                976
                      1163
                                 RETICULON.
FT
    DOMAIN
                  33
                        46
                                 POLY-GLU.
FT
    DOMAIN
                 73
                        76
                                 POLY-ALA.
FT
    DOMAIN
                140
                       145
                                 POLY-PRO.
FT
    VARSPLIC
                 1
                       964
                                 Missing (in isoform 3).
FT
                                 /FTId=VSP 005656.
FT
    VARSPLIC
                965
                       975
                                 AVLSAELSKTS -> MDGQKKHWKDK (in isoform
FT
                                 3).
```

```
FT
                           /FTId=VSP 005657.
FT
    VARSPLIC
              173
                   975
                           Missing (in isoform 2).
FT
                           /FTId=VSP 005658.
FT
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             192
                   975
                           Missing (in isoform 4).
FT
                           /FTId=VSP 005659.
FT
    CONFLICT
             1130
                  1131
                           MISSING (IN REF. 3; AAD31020).
SO
    SEOUENCE
             1163 AA;
                    126386 MW; 8CB894B09E94F0B6 CRC64;
  Query Match
                    100.0%;
                           Score 5848; DB 1; Length 1163;
  Best Local Similarity
                    100.0%; Pred. No. 9.1e-214;
  Matches 1163; Conservative
                          0;
                            Mismatches
                                        0;
                                           Indels
                                                   0;
                                                      Gaps
                                                            0;
Qу
          1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
           ]]]]]]]
Db
         1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
         61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAAPAPSLP 120
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Db
        121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
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           Db
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           241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300
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           Db
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Qу
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Db
        901 DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED 960
Qу
           901 DEVHVSDEFSENRSSVSKASISPSNVSALEPOTEMGSIVKSKSLTKEAEKKLPSDTEKED 960
Db
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Qу
           961 RSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLS 1020
Db
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QУ
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Db
       1081 FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHOVOIDHYLGLA 1140
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Db
Qу
       1141 NKSVKDAMAKIQAKIPGLKRKAD 1163
           Dħ
       1141 NKSVKDAMAKIQAKIPGLKRKAD 1163
RESULT 2
RTN4 HUMAN
ID
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AC
AC
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DT
   28-FEB-2003 (Rel. 41, Created)
DT
    28-FEB-2003 (Rel. 41, Last sequence update)
DT
    15-SEP-2003 (Rel. 42, Last annotation update)
   Reticulon 4 (Neurite outgrowth inhibitor) (Nogo protein) (Foocen)
DE
DE
    (Neuroendocrine-specific protein) (NSP) (Neuroendocrine specific
   protein C homolog) (RTN-x) (Reticulon 5) (My043 protein).
DE
GN
   RTN4 OR NOGO OR ASY OR KIAA0886.
```

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Prinjha R., Moorė S.E., Vinson M., Blake S., Morrow R., Christie G.,

Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

SEQUENCE FROM N.A. (ISOFORMS 1; 2 AND 3).

Michalovich D., Simmons D.L., Walsh F.S.;

MEDLINE=20129242; PubMed=10667780;

OS

OC

OC

OX

RN RP

RX

RA

RA

Homo sapiens (Human).

NCBI TaxID=9606;

```
RT
      "Inhibitor of neurite outgrowth in humans.";
 RL
      Nature 403:383-384(2000).
RN
RP
      SEQUENCE FROM N.A. (ISOFORMS 1 AND 2).
RC
      TISSUE=Brain;
      MEDLINE=21010696; PubMed=11126360;
RX
      Tagami S., Eguchi Y., Kinoshita M., Takeda M., Tsujimoto Y.;
RA
RT
      "A novel protein, RTN-XS, interacts with both Bcl-XL and Bcl-2 on
      endoplasmic reticulum and reduces their anti-apoptotic activity.";
RT
RL
      Oncogene 19:5736-5746(2000).
RN
      [3]
RΡ
      SEQUENCE FROM N.A. (ISOFORMS 1; 2 AND 3).
RX
      MEDLINE=20237542; PubMed=10773680;
RA
      Yang J., Yu L., Bi A.D., Zhao S.-Y.;
RT
      "Assignment of the human reticulon 4 gene (RTN4) to chromosome
      2p14-->2p13 by radiation hybrid mapping.";
RT
RL
     Cytogenet. Cell Genet. 88:101-102(2000).
RN
      [4]
RP
     SEQUENCE FROM N.A. (ISOFORM 4).
RA
     Jin W.-L., Ju G.;
      "Developmentally-regulated alternative splicing in a novel Nogo-A.";
     Submitted (NOV-2000) to the EMBL/GenBank/DDBJ databases.
RI.
RN
RP
     SEQUENCE FROM N.A. (ISOFORMS 2 AND 3).
RC
     TISSUE=Placenta, and Skeletal muscle;
RA
     Ito T., Schwartz S.M.;
RT
     "Cloning of a member of the reticulon gene family in human.";
     Submitted (FEB-1999) to the EMBL/GenBank/DDBJ databases.
RL
RN
RΡ
     SEQUENCE FROM N.A. (ISOFORM 2).
RC
     TISSUE=Fibroblast:
     Yutsudo M.;
RA
RT
     "Isolation of a cell death-inducing gene.";
RL
     Submitted (JUN-1998) to the EMBL/GenBank/DDBJ databases.
RN
RΡ
     SEQUENCE FROM N.A. (ISOFORM 3).
RC
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     Song H., Peng Y., Zhou J., Huang Q., Dai M., Mao Y.M., Yu Y., Xu X.,
RA
     Luo B., Hu R., Chen J.;
RA
RT
     "Human neuroendocrine-specific protein C (NSP) homolog gene.";
RL
     Submitted (JUL-1998) to the EMBL/GenBank/DDBJ databases.
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     SEQUENCE FROM N.A. (ISOFORM 3).
     Gu J.R., Wan D.F., Zhao X.T., Zhou X.M., Jiang H.Q., Zhang P.P.,
RA
     Qin W.X., Huang Y., Qiu X.K., Qian L.F., He L.P., Li H.N., Yu Y.,
RA
RA
     Yu J., Han L.H.;
RT
     "Novel human cDNA clone with function of inhibiting cancer cell
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     growth.";
     Submitted (AUG-1999) to the EMBL/GenBank/DDBJ databases.
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     SEQUENCE FROM N.A. (ISOFORM 1).
RC
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     Nagase T., Ishikawa K.-I., Suyama M., Kikuno R., Hirosawa M.,
RA
     Miyajima N., Tanaka A., Kotani H., Nomura N., Ohara O.;
RT
     "Prediction of the coding sequences of unidentified human genes. XII.
     The complete sequences of 100 new cDNA clones from brain which code
RT
```

```
RT
     for large proteins in vitro.";
RL
     DNA Res. 5:355-364(1998).
RN
RP
     SEQUENCE FROM N.A. (ISOFORMS 2 AND 3).
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     TISSUE=Brain, Pancreas, Placenta, and Skeletal muscle;
RX
     MEDLINE=22388257; PubMed=12477932;
     Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RA
     Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
RA
RA
     Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
     Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA
RA
     Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
     Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA
RA
     Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RA
     Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA
     Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
     Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RΑ
     Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
RA
     Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,
RA
RA
     Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA
     Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
     Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA
RA
     Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
     Schnerch A., Schein J.E., Jones S.J.M., Marra M.A.;
RA
     "Generation and initial analysis of more than 15,000 full-length
RT
     human and mouse cDNA sequences.";
RT
RL
     Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
RN
RΡ
     SEQUENCE FROM N.A. (ISOFORM 3).
     MEDLINE=20499367; PubMed=11042152;
RX
     Zhang Q.-H., Ye M., Wu X.-Y., Ren S.-X., Zhao M., Zhao C.-J., Fu G.,
RA
RA
     Shen Y., Fan H.-Y., Lu G., Zhong M., Xu X.-R., Han Z.-G., Zhang J.-W.,
     Tao J., Huang Q.-H., Zhou J., Hu G.-X., Gu J., Chen S.-J., Chen Z.;
RA
     "Cloning and functional analysis of cDNAs with open reading frames for
RT
RT
     300 previously undefined genes expressed in CD34+ hematopoietic
RT
     stem/progenitor cells.";
RL
     Genome Res. 10:1546-1560(2000).
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     SEQUENCE OF 482-1192 FROM N.A. (ISOFORM 1/4).
RC
     TISSUE=Brain;
RA
     Mao Y.M., Xie Y., Zheng Z.H.;
RL
     Submitted (MAY-1998) to the EMBL/GenBank/DDBJ databases.
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RC
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     Sha J.H., Zhou Z.M., Li J.M.;
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RL
     Submitted (JAN-2001) to the EMBL/GenBank/DDBJ databases.
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RC
     TISSUE=Brain;
RX
     MEDLINE=20129259; PubMed=10667797;
     GrandPre T., Nakamura F., Vartanian T., Strittmatter S.M.;
RT
     "Identification of the Nogo inhibitor of axon regeneration as a
RT
     Reticulon protein.";
RL
     Nature 403:439-444(2000).
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     [15]
RP
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     TISSUE=Brain;
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```
RX
     MEDLINE=21069055; PubMed=11201742;
RA
     Fournier A.E., Grandpre T., Strittmatter S.M.;
     "Identification of a receptor mediating Nogo-66 inhibition of axonal
RT
RT
     regeneration.";
RL
     Nature 409:341-346(2001).
RN
RP
     REVIEW.
RX
     MEDLINE=21888956; PubMed=11891768;
RΑ
     Ng C.E.L., Tang B.L.;
     "Nogos and the Nogo-66 receptor: factors inhibiting CNS neuron
RT
RT
     regeneration.";
RL
     J. Neurosci. Res. 67:559-565(2002).
CC
     -!- FUNCTION: Potent neurite outgrowth inhibitor which may also help
CC
         block the regeneration of the nervous central system in adults.
CC
         Isoform 2 reduces the anti-apoptotic activity of Bcl-xl and Bcl-2.
CC
         This is likely consecutive to their change in subcellular
CC
         location, from the mitochondria to the endoplasmic reticulum,
CC
         after binding and sequestration.
CC
     -!- SUBUNIT: Binds to RTN4R. Interacts with Bcl-xl and Bcl-2.
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein. Endoplasmic
CC
         reticulum. Anchored to the membrane of the endoplasmic reticulum
CC
         through 2 putative transmembrane domains.
CC
     -! - ALTERNATIVE PRODUCTS:
CC
         Event=Alternative splicing; Named isoforms=4;
CC
         Name=1; Synonyms=RTN 4A, Nogo-A, RTN-xL;
CC
           IsoId=Q9NQC3-1; Sequence=Displayed;
CC
         Name=2; Synonyms=RTN 4B, Nogo-B, RTN-xS, Foocen-M;
CC
           IsoId=Q9NQC3-2; Sequence=VSP 005655;
CC
         Name=3; Synonyms=RTN 4C, Nogo-C, Foocen-S;
CC
           IsoId=Q9NQC3-3; Sequence=VSP 005652, VSP 005653;
CC
         Name=4;
CC
           IsoId=Q9NQC3-4; Sequence=VSP 005654;
CC
     -!- TISSUE SPECIFICITY: Isoform 1 is specifically expressed in brain
CC
         and testis and weakly in heart and skeletal muscle. Isoform 2 is
CC
         widely expressed excepted for the liver. Isoform 3 is expressed in
CC
         brain, skeletal muscle and adipocytes. Isoform 4 is testis-
CC
         specific.
CC
     -!- SIMILARITY: Contains 1 reticulon domain.
     -!- CAUTION: Ref.11 sequence differs from that shown due to
CC
         frameshifts in positions 1149 and 1156.
CC
     _____
CC
     This SWISS-PROT entry is copyright. It is produced through a collaboration
CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
CC
     the European Bioinformatics Institute. There are no restrictions on its
CC
     use by non-profit institutions as long as its content is in no way
CC
     modified and this statement is not removed. Usage by and for commercial
CC
     entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
     or send an email to license@isb-sib.ch).
CC
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DR
     EMBL; AJ251384; CAB99249.1; -.
DR
     EMBL; AJ251385; CAB99250.1; -.
DR
     EMBL; AB040462; BAB18927.1; -.
DR
     EMBL; AB040463; BAB18928.1; -.
DR
     EMBL; AF148537; AAG12176.1; -.
DR
     EMBL; AF148538; AAG12177.1; -.
DR
     EMBL; AF087901; AAG12205.1; -.
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DR
   EMBL; AF320999; AAG40878.1; -.
   EMBL; AF132047; AAD31021.1; -.
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   EMBL; AF132048; AAD31022.1;
DR
   EMBL; AB015639; BAA83712.1; -.
DR
DR
   EMBL; AF077050; AAD27783.1; -.
DR
   EMBL; AF177332; AAG17976.1; -.
   EMBL; AB020693; BAA74909.1; -.
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    EMBL; BC001035; AAH01035.1; -.
DR
DR
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QУ
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Db
Qу
        61 PAAGLSAAAVP--PAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWERSPAA---P 115
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        59 PAAGLSAAPVPTAPAAGAPLMDFGNDFVPPAPRGPLPAAPPVAPEROPSWDPSPVSSTVP 118
        116 APSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAE----
                                               -PAAPPSTPAAPKR 166
Qу
               119 APSPLSAAAVSPSKLPEDDEPPARPPPPPPASVSPQAEPVWTPPAPAPAAPPSTPAAPKR 178
Db
        167 RG-SGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPS 225
Qy
           179 RGSSGSVDETLFALPAASEPVIRSSAEN-MDLKEOPGNTISAGOEDFPSVLLETAASLPS 237
Db
       226 LSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEY 285
QУ
                238 LSPLSAASFKEHEYLGNLSTVLPTEGTLQENVSEASKEVSEKAKTLLIDRDLTEFSELEY 297
Db
       286 SEMGSSFKGSPKGESAILVENTKEEVIVRSKDKED-LVCSAALHSPQESPVG-----KED 339
QУ
                 Db
       298 SEMGSSFSVSPKAESAVIVANPREEIIVKNKDEEEKLVSNNILHNOOELPTALTKLVKED 357
       340 RVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAA----RANV 395
Qу
            358 EVVSSEKAKDSFNEKRVAVEAPMREEYADFKPFERVWEVKDSKEDS-DMLAAGGKIESNL 416
Db
       396 ESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFT-SATESTT 454
Qу
           Db
       417 ESKVDKKCFADSLEQTNHEKDSESSNDDTSFPSTPEGIKDRSGAYITCAPFNPAATESIA 476
Qу
       455 ANTFPLLEDHTSENKTDEKKIEERKAQIITEK-TSPKTSNPFLVAVQDSEADYVTTDTLS 513
            477 TNIFPLLGDPTSENKTDEKKIEEKKAQIVTEKNTSTKTSNPFLVAAQDSETDYVTTDNLT 536
Db
       514 KVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQL 573
Qу
               537 KVTEEVVANMPEGLTPDLVQEACESELNEVTGTKIAYETKMDLVQTSEVMQESLYPAAQL 596
Db
       574 CPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPEN 633
Qу
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Db
        597 CPSFEESEATPSPVLPDIVMEAPLNSAVPSAGASVIQPSSSPLEA-SSVNYESIKHEPEN 655
Qу
        634 PPPYEEAMNVALKAL-GTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDF 692
            656 PPPYEEAMSVSLKKVSGIKEEIKEPENINAALQETEAPYISIACDLIKETKLSAEPAPDF 715
Db
        693 SNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVS 752
Qу
            716 SDYSEMAKVEQPVPDHSELVEDSSPDSEPVDLFSDDSIPDVPQKQDETVMLVKESLTETS 775
Db
        753 -ETVAQHK-EERLSASPQELGKPYLESFQPNLHSTKDA-ASNDIPTLTKKEKISLQMEEF 809
Qу
            776 FESMIEYENKEKLSALPPEGGKPYLESFKLSLDNTKDTLLPDEVSTLSKKEKIPLQMEEL 835
Db
        810 NTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDS-PKLAKEYTDLEVS 868
Qу
            836 STAVYSNDDLFISKEAQIRETETFSDSSPIEIIDEFPTLISSKTDSFSKLAREYTDLEVS 895
Db
        869 DKSEIANIQSGADSLPCLELPCDLSFKNIYPK--DEVHVSDEFSENRSSVSKASISPSNV 926
Qу
                   896 HKSEIANAPDGAGSLPCTELPHDLSLKNIQPKVEEKISFSDDFSKNGSATSKVLLLPPDV 955
Db
        927 SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIK 986
Qу
               Db
        956 SALATQAEIESIVKPKVLVKEAEKKLPSDTEKEDRSPSAIFSAELSKTSVVDLLYWRDIK 1015
        987 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA 1046
Qу
           Db
       1016 KTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRA 1075
       1047 YLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1106
Qу
           1076 YLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALF 1135
Dh
Qу
       1107 NGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
           1136 NGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 1192
Db
RESULT 3
RTN4 MOUSE
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ID
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AC
    28-FEB-2003 (Rel. 41, Created)
DT
DT
    28-FEB-2003 (Rel. 41, Last sequence update)
    28-FEB-2003 (Rel. 41, Last annotation update)
DT
    Reticulon 4 (Neurite outgrowth inhibitor) (Nogo protein).
DE
GN
    RTN4 OR NOGO.
OS
    Mus musculus (Mouse).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
0^{\circ}
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OC
OX
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RN
RΡ
    SÈQUENCE FROM N.A.
RC
    STRAIN=3T3-L1; TISSUE=Adipocyte;
RA
    Coulson A.C., Craggs P.D., Morris N.J.;
```

"Mouse vp20/RTN4C cDNA.";

RT

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RL
     Submitted (DEC-2000) to the EMBL/GenBank/DDBJ databases.
RN
     SEQUENCE OF 170-199 FROM N.A.
RP
RC
     STRAIN=C57BL/6J; TISSUE=Embryo;
RX
     MEDLINE=21085660; PubMed=11217851;
RA
     Kawai J., Shinagawa A., Shibata K., Yoshino M., Itoh M., Ishii Y.,
RA
     Arakawa T., Hara A., Fukunishi Y., Konno H., Adachi J., Fukuda S.,
     Aizawa K., Izawa M., Nishi K., Kiyosawa H., Kondo S., Yamanaka I.,
RA
     Saito T., Okazaki Y., Gojobori T., Bono H., Kasukawa T., Saito R.,
RA
RA
     Kadota K., Matsuda H.A., Ashburner M., Batalov S., Casavant T.,
     Fleischmann W., Gaasterland T., Gissi C., King B., Kochiwa H.,
RA
RA
     Kuehl P., Lewis S., Matsuo Y., Nikaido I., Pesole G., Quackenbush J.,
     Schriml L.M., Staubli F., Suzuki R., Tomita M., Wagner L., Washio T.,
RA
     Sakai K., Okido T., Furuno M., Aono H., Baldarelli R., Barsh G.,
RA
RA
     Blake J., Boffelli D., Bojunga N., Carninci P., de Bonaldo M.F.,
RA
     Brownstein M.J., Bult C., Fletcher C., Fujita M., Gariboldi M.,
     Gustincich S., Hill D., Hofmann M., Hume D.A., Kamiya M., Lee N.H.,
RA
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     Lyons P., Marchionni L., Mashima J., Mazzarelli J., Mombaerts P.,
RA
     Nordone P., Ring B., Ringwald M., Rodriguez I., Sakamoto N.,
     Sasaki H., Sato K., Schoenbach C., Seya T., Shibata Y., Storch K.-F.,
RA
RA.
     Suzuki H., Toyo-oka K., Wang K.H., Weitz C., Whittaker C., Wilming L.,
     Wynshaw-Boris A., Yoshida K., Hasegawa Y., Kawaji H., Kohtsuki S.,
RA
RA
     Hayashizaki Y.;
     "Functional annotation of a full-length mouse cDNA collection.";
RT
RL
     Nature 409:685-690(2001).
     -!- FUNCTION: Potent neurite outgrowth inhibitor which may also help
CC
CC
         block the regeneration of the nervous central system in adults (By
CC
         similarity).
CC
     -!- SUBUNIT: Binds to RTN4R. Interacts with Bcl-xl and Bcl-2 (By
CC
         similarity).
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein. Anchored to the
         membrane of the endoplasmic reticulum through 2 putative
CC
CC
         transmembrane domains (By similarity).
CC
     -!- ALTERNATIVE PRODUCTS:
CC
         Event=Alternative splicing; Named isoforms=1;
CC
           Comment=A number of isoforms may be produced:
CC
         Name=1;
CC
           IsoId=Q99P72-1; Sequence=Displayed;
CC
     -!- SIMILARITY: Contains 1 reticulon domain.
CC
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CC
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     or send an email to license@isb-sib.ch).
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     EMBL; AF326337; AAK08076.1; -.
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     EMBL; AK003859; -; NOT ANNOTATED CDS.
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    MGD; MGI:1915835; Rtn4.
    GO; GO:0030176; C:endoplasmic reticulum membrane, intrinsic p. . .; ISS.
DR
DR
    GO; GO:0005783; C:endoplasmic reticulum; IDA.
DR
    GO; GO:0005635; C:nuclear membrane; ISS.
DR
    GO; GO:0005515; F:protein binding activity; ISS.
DR
    GO; GO:0019987; P:negative regulation of anti-apoptosis; ISS.
    GO; GO:0030517; P:negative regulation of axon extension; ISS.
DR
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InterPro; IPR003388; Reticulon.
DR
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     PROSITE; PS50845; RETICULON; 1.
DR
     Endoplasmic reticulum; Alternative splicing; Transmembrane.
KW
FΤ
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                                CYTOPLASMIC (Potential).
                 1
                        25
FT
     TRANSMEM
                 26
                        50
                                POTENTIAL.
FT
     DOMAIN
                 51
                       137
                                LUMENAL (Potential).
FT
     TRANSMEM
                138
                       162
                                POTENTIAL.
FT
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                163
                       199
                                CYTOPLASMIC (Potential).
FT.
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                 12
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                               1; Mismatches
                                                3; Indels
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          973 KTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVI 1032
Qу
                Db
           9 KDKVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVI 68
         1033 QAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKF 1092
Qу
             Db
          69 QAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKF 128
         1093 AVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQ 1152
Οv
             Db
         129 AVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQAQIDHYLGLANKSVKDAMAKIQ 188
Qу
        1153 AKIPGLKRKAD 1163
             Db
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RESULT 4
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                                 PRT;
                                       777 AA.
AC
     Q64548; Q64547;
     16-OCT-2001 (Rel. 40, Created)
DT
DT
     16-OCT-2001 (Rel. 40, Last sequence update)
DT
     15-SEP-2003 (Rel. 42, Last annotation update)
DE
    Reticulon 1 (Neuroendocrine-specific protein) (S-rex).
GN
    RTN1 OR NSP.
    Rattus norvegicus (Rat).
OS
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX
    NCBI TaxID=10116;
RN
    [1]
RΡ
    SEQUENCE FROM N.A. (ISOFORMS RTN1-B AND RTN1-S).
RC
    STRAIN=Wistar; TISSUE=Brain cortex;
    MEDLINE=96386034; PubMed=8793864;
RX
RA
    Baka I.D., Ninkina N.N., Pinon L.G.P., Adu J., Davies A.M.,
RA
    Georgiev G.P., Buchman V.L.;
RT
    "Intracellular compartmentalization of two differentially spliced s-
    rex/NSP mRNAs in neurons.";
RT
RL
    Mol. Cell. Neurosci. 7:289-303(1996).
CC
    -!- FUNCTION: MAY BE INVOLVED IN NEUROENDOCRINE SECRETION OR IN
CC
        MEMBRANE TRAFFICKING IN NEUROENDOCRINE CELLS.
```

DR

```
CC
     -!- SUBCELLULAR LOCATION: ENDOPLASMIC RETICULUM MEMBRANE (BY
 CC
         SIMILARITY).
 CC
     -!- ALTERNATIVE PRODUCTS:
 CC
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         Name=RTN1-B; Synonyms=S-RexB;
 CC
 CC
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 CC
         Name=RTN1-S; Synonyms=S-RexS;
CC
           IsoId=Q64548-2; Sequence=VSP 005647, VSP 005648;
     -!- TISSUE SPECIFICITY: EXPRESSED PREDOMINANTLY IN CENTRAL AND
CC
         PERIPHERAL NERVOUS SYSTEM OF NEWBORN AND ADULT RATS. LOW LEVELS
CC
CC
        HAVE BEEN ALSO DETECTED IN HEART, ADRENAL GLAND AND SPLEEN.
        EXPRESSION OF ISOFORM RTN1-B IS RESTRICTED TO PARTICULAR NEURONAL
CC
CC
        TYPES.
CC
     -!- DEVELOPMENTAL STAGE: DETECTED ON EMBRYONIC DAY E10 IN THE
CC
        HINDBRAIN AND IN E11 IN THE FOREBRAIN. DURING THE NEXT 3 EMBRYONIC
CC
        DAYS THE LEVELS OF S-REXS INCREASES AND REMAINS STABLE AT E13 IN
        THE HINDBRAIN AND AT E14 IN THE FOREBRAIN. THE LEVELS OF S-REXB
CC
CC
        DOES NOT CHANGE AS SIGNIFICANTLY DURING DEVELOPMENT OF THE
CC
        HINDBRAIN.
CC
     -!- SIMILARITY: Contains 1 reticulon domain.
CC
     ______
CC
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CC
     between the Swiss Institute of Bioinformatics and the EMBL outstation -
     the European Bioinformatics Institute. There are no restrictions on its
CC
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     entities requires a license agreement (See http://www.isb-sib.ch/announce/
CC
     or send an email to license@isb-sib.ch).
CC
     CC
DR
     EMBL; U17604; AAC53046.1; -.
     EMBL; U17603; AAC53045.1; -.
DR
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     InterPro; IPR003388; Reticulon.
DR
     Pfam; PF02453; Reticulon; 1.
     PROSITE; PS50845; RETICULON; 1.
DR
KW
     Endoplasmic reticulum; Alternative splicing; Transmembrane.
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FT
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FT
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FT
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  Matches 258; Conservative 109; Mismatches 264; Indels 166; Gaps
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         485 EKTSPKTSNPFLVAVQDSE-----ADYVTTDTL---SKVTEAAVSNMPEGL--TPD 530
             29 EEATPKGARP---AQQDGEPAWGSGAGAGVVSSRGLCSGPARSPPVAMETASTGVAAVPD 85
Db
         531 LVQEACESELNEATG-----TKIAYETKVDLVQTSEAIQ-ESLYPTAQLCPSFEEAEA 582
Qу
              Db
         86 ALDHSSSPTLKDGEGACYTSLISDICYPPREDSAYFTGILQKENGHITTSESP---EELG 142
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Qy		TPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLE	
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Qу	631	PENPPPYEEAMNVALKALGTK-EGIKEPESFNAAVQETE	668
Db	199	RPQEAKGQEEQSPGLEDKDLDFKDKDSEVSTKPEGVHAPNQPSPVEGKLIKDNLFEESTF	258
QУ	669	APYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVD	723
Db	259	APYIDELSDEQHRMSLVTAPVKITLTEIGPPVMTATHETIPEKQDLCLKPSPDTVPTV	316
Qy	724	LFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESF	778
Db	317	: : : : : : : : : : : :	370
Qу	779	QPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETF	833
Db	371	: : :	414
Qу	834	SDSSPIEIIDEFPTFVSAKDDSPKLAK-EYTDLEVSDKSEIANIQSGADSLPCL	886
Db	415	: : : : : : :: : :	466
QУ	887	ELPCDL-SFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEM	935
Db	467		525
Qу	936	KSKSLTKEAEKKLPSDTEKEDRSLSAVLSAEL	971
Db	526	: : : : PVTLQSRPEPSSGDGAPVPEPPKSQQQKPEEEAVSSSQSPAATEIPGPLGSDLVP	580
Qу	972	SKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFR: : : : :	1026
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Db	641	: : : : : : : :: : :	700
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Db	701	: :: :: : : : : : :	760
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Db	761	: : VVAKIQAKIPGAKRHAE 777	
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DE
    Reticulon 1 (Neuroendocrine-specific protein).
GN
    RTN1 OR NSP.
OS
    Homo sapiens (Human).
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
OX
    NCBI TaxID=9606;
RN
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    TISSUE=Lung carcinoma;
    MEDLINE=93293865; PubMed=7685762;
RX
    Roebroek A.J.M., Van de Velde H.J.K., Van Bokhoven A., Broers J.L.V.,
RA
    Ramaekers F.C.S., Van de Ven W.J.M.;
RA
     "Cloning and expression of alternative transcripts of a novel
RT
    neuroendocrine-specific gene and identification of its 135-kDa
RT
RT
    translational product.";
    J. Biol. Chem. 268:13439-13447(1993).
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    ALTERNATIVE SPLICING.
RP
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    MEDLINE=96429995; PubMed=8833145;
RA
     Roebroek A.J.M., Ayoubi T.A.Y., Van de Velde H.J.K.,
RA
     Schoenmakers E.F.P.M., Pauli I.G.L., Van de Ven W.J.M.;
RT
    "Genomic organization of the human NSP gene, prototype of a novel gene
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     family encoding reticulons.";
RL
    Genomics 32:191-199(1996).
RN
RΡ
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    MEDLINE=98228245; PubMed=9560466;
     Hens J., Nuydens R., Geerts H., Senden N.H., Van de Ven W.J.M.,
RA
RA
     Roebroek A.J., van de Velde H.J., Ramaekers F.C., Broers J.L.;
     "Neuronal differentiation is accompanied by NSP-C expression.";
RT
RL
     Cell Tissue Res. 292:229-237(1998).
CC
     -!- FUNCTION: MAY BE INVOLVED IN NEUROENDOCRINE SECRETION OR IN
CC
        MEMBRANE TRAFFICKING IN NEUROENDOCRINE CELLS.
CC
     -!- SUBCELLULAR LOCATION: Endoplasmic reticulum membrane.
CC
     -!- ALTERNATIVE PRODUCTS:
CC
        Event=Alternative splicing; Named isoforms=3;
CC
        Name=RTN1-A; Synonyms=NSP-A;
CC
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CC
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CC
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CC
     -!- TISSUE SPECIFICITY: EXPRESSED IN NEURAL AND NEUROENDOCRINE TISSUES
CC
        AND CELL CULTURES DERIVED THEREFROM. EXPRESSION OF ISOFORM RTN1-C
CC
        IS STRONGLY CORRELATED WITH NEURONAL DIFFERENTIATION.
CC
     -!- PTM: ISOFORMS RTN1-A AND RTN1-B ARE PHOSPHORYLATED.
CC
     -!- SIMILARITY: Contains 1 reticulon domain.
CC
     ______
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     or send an email to license@isb-sib.ch).
CC
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    PIR; A46583; A46583.
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DR
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DR
    MIM; 600865; -.
    GO; GO:0030176; C:endoplasmic reticulum membrane, intrinsic p. . .; TAS.
DR
    GO; GO:0004871; F:signal transducer activity; NAS.
DR
    GO; GO:0030182; P:neuron differentiation; TAS.
DR
    GO; GO:0007165; P:signal transduction; NAS.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam: PF02453; Reticulon; 1.
DR
DR
    PROSITE; PS50845; RETICULON; 1.
    Endoplasmic reticulum; Alternative splicing; Transmembrane;
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                      623
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FΤ
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FT
    TRANSMEM
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                               RETICULON.
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FT
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FT
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          65 SGPARQSP--VAMETASTGVAGVSSAMDHTFSTTSKDGEG-----SCYTSLI----S 110
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         547 KIAYETKVDLVQTSEAIQ-ESLYPTAQLCPSFEEAEATPSPVLPDI--VMEAPLNS---- 599
Qу
              111 DICYPPQEDSTYFTGILQKENGHVTISESP---EELGTPGPSLPDVPGIESRGLFSSDSG 167
Dh
         600 --LLPSAGASVVQPSVSPLEAPPPVSY------DSIKLEPENPPPYEEA------M 641
Qу
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         168 IEMTPAESTEVNKILADPLDQMKAEAYKYIDITRPEEVKHQEQHHPELEDKDLDFKNKDT 227
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Qу
                                       : : | | | | |
                                                      ::::
                     ||::||:
         228 DISIK----PEGVREPDKPAPVEGKIIKDHLLEESTFAPYID---DLSEEQRRAPQITTP 280
Db
         691 DFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLF-SDDSIPEV----PQTQEEAVMLMK 745
Qу
                              :: :|| : : | | |::| | |::
                  : | |
         281 VKITLTEIE-----PSVETTTQEKTPEKQDICLKPSPDTVPTVTVSEPEDDSPGSITPP 334
Db
         746 ESLTEVSETVAQHK----EERLSASPQELGKPYLESFQP----NLHSTKDAASND---- 792
Qу
              335 SSGTEPSAAESQGKGSISEDELITAIKEAKGLSYETAENPRPVGQLADRPEVKARSGPPT 394
Db
         793 IPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKI----- 827
QУ
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Db
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Qу
               : | | | | | | | | |
                                  453 ILREEREAELDS---ELIIESCDASSASEESPKREODSPPMKPSALDAIREETGVRAEER 509
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        880 ADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPOTEMGSIV 939
QУ
                Db
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        940 KSKSLTKEAEKKLPSDTEKEDRSLSAVLSAE-----LSKTSVVDLLYWRD 984
Ov
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        550 -----LPRKPEEDSSSNQSPAATKGPGPLGPGAPPPLLFLNKQKAIDLLYWRD 597
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            598 IKQTGIVFGSFLLLLFSLTQFSVVSVVAYLALAALSATISFRIYKSVLQAVQKTDEGHPF 657
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Db
        1105 LFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
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Db
RESULT 6
RTN3 HUMAN
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AC
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DT
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    16-OCT-2001 (Rel. 40, Last sequence update)
דת
DТ
    15-SEP-2003 (Rel. 42, Last annotation update)
DE
    Reticulon protein 3 (Neuroendocrine-specific protein-like 2) (NSP-like
DE
    protein II) (NSPLII).
GN
    RTN3 OR NSPL2.
OS
    Homo sapiens (Human).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
    NCBI TaxID=9606;
OX
RN
    [1]
RP ·
    SEQUENCE FROM N.A., AND TISSUE SPECIFICITY.
RC
    TISSUE=Retina;
    MEDLINE=99265974; PubMed=10331947;
RX
    Moreira E.F., Jaworski C.J., Rodriguez I.R.;
RA
RT
    "Cloning of a novel member of the reticulon gene family (RTN3): gene
RT
    structure and chromosomal localization to 11q13.";
RL
    Genomics 58:73-81(1999).
RN
RP
    SEQUENCE FROM N.A.
RA
    Huang X., Zhou Y., Du G., Yuan J., Qiang B.;
RТ
    "Cloning and expression analysis of a cDNA encoding a novel
RT
    neuroendocrine-specific protein-like protein 1: NSPL1.";
RL
    Submitted (JAN-1999) to the EMBL/GenBank/DDBJ databases.
RN
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RP
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     TISSUE=Brain, Eye, and Lymph;
RX
     MEDLINE=22388257; PubMed=12477932;
     Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RA
RA
     Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
     Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
RA
RA
     Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
RA
     Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
     Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA
     Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
RA
     Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
     Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
RΆ
RA
     Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA
     Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
RΑ
     Fahey J., Helton E., Ketteman M., Madan A., Rodriques S., Sanchez A.,
     Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA
     Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA
     Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA
     Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
RA
     Schnerch A., Schein J.E., Jones S.J.M., Marra M.A.;
RT
     "Generation and initial analysis of more than 15,000 full-length
RT
     human and mouse cDNA sequences.";
RL
     Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
CC
     -!- SUBCELLULAR LOCATION: Integral membrane protein. Endoplasmic
CC
         reticulum (Potential).
CC
     -!- TISSUE SPECIFICITY: WIDELY EXPRESSED WITH HIGHEST EXPRESSION IN
CC
         BRAIN. THREE TIMES MORE ABUNDANT IN MACULA THAN IN PERIPHERAL
CC
         RETINA.
CC
     -!- SIMILARITY: Contains 1 reticulon domain.
CC
     ______
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CC
DR
     EMBL; AF059524; AAC99319.1; -.
DR
     EMBL; AF059529; AAD20951.1; -.
     EMBL; AF059525; AAD20951.1; JOINED.
DR
DR
     EMBL; AF059526; AAD20951.1; JOINED.
DR
     EMBL; AF059527; AAD20951.1; JOINED.
DR
     EMBL; AF059528; AAD20951.1; JOINED.
DR
     EMBL; AF119297; AAD26810.1; -.
DR
     EMBL; BC000634; AAH00634.1; -.
     EMBL; BC010556; AAH10556.1; -.
DR
     EMBL; BC011394; AAH11394.1; -.
DR
DR
     EMBL; BC022993; AAH22993.1; -.
DR
     Genew; HGNC:10469; RTN3.
DR
    MIM; 604249; -.
DR
     GO; GO:0005615; C:extracellular space; TAS.
DR
     InterPro; IPR003388; Reticulon.
DR
     Pfam; PF02453; Reticulon; 1.
DR
     PROSITE; PS50845; RETICULON; 1.
ΚW
     Transmembrane; Endoplasmic reticulum.
FT
     TRANSMEM
                                 POTENTIAL.
              68
                        88
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FT
    TRANSMEM
              177 197
                             POTENTIAL.
FT
    DOMAIN
              48
                    236
                             RETICULON.
SO
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             236 AA; 25609 MW; DDC6A4544ABCDFB7 CRC64;
                      10.7%; Score 625.5; DB 1; Length 236;
 Best Local Similarity 59.1%; Pred. No. 1.4e-17;
 Matches 114; Conservative 41; Mismatches 37; Indels
                                                       1; Gaps
        972 SKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGV 1031
Qу
            Db
         44 SSCAVHDLIFWRDVKKTGFVFGTTLIMLLSLAAFSVISVVSYLILALLSVTISFRIYKSV 103
Оv
        1032 IQAIQKSDEGHPFRAYLESEVAISEELVOKYSNSALGHVNSTIKELRRLFLVDDLVDSLK 1091
            Db
        104 IQAVQKSEEGHPFKAYLDVDITLSSEAFHNYMNAAMVHINRALKLIIRLFLVEDLVDSLK 163
       1092 FAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKI 1151
QУ
             | : ||
        164 LAVFMWLMTYVGAVFNGITLLILAELLIFSVPIVYEKYKTQIDHYVGIARDQTKSIVEKI 223
Db
Qу
       1152 QAKIPGL-KRKAD 1163
            |||:||: |:||:
Dh
        224 QAKLPGIAKKKAE 236
RESULT 7
RTN3 MOUSE
ID
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                 STANDARD;
                              PRT;
                                    237 AA.
AC
    Q9ES97;
DT
    16-OCT-2001 (Rel. 40, Created)
DT
    16-OCT-2001 (Rel. 40, Last sequence update)
    28-FEB-2003 (Rel. 41, Last annotation update)
DΕ
    Reticulon protein 3.
GN
    RTN3.
OS
    Mus musculus (Mouse).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC.
OC
    Mammalia; Eutheria; Rodentia; Sciuroqnathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
    [1]
RP
    SEQUENCE FROM N.A.
RA
    Huang X., Zhou Y., Qiang H., Yuan J., Qiang B.;
RT
    "Cloning and expression profile of a novel mouse cDNA encoding a human
RT
    RTN3 homolog.";
RL
    Submitted (OCT-1999) to the EMBL/GenBank/DDBJ databases.
CC
    -!- SUBCELLULAR LOCATION: Integral membrane protein. Endoplasmic
CC
       reticulum (Potential).
CC
    -!- SIMILARITY: Contains 1 reticulon domain.
CC
    CC
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    CC
DR
    EMBL; AF195940; AAG31360.1; -.
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DR
    MGD; MGI:1339970; Rtn3.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
DR.
    PROSITE; PS50845; RETICULON; 1.
KW
    Transmembrane; Endoplasmic reticulum.
FT
    TRANSMEM
                69
                       89
                               POTENTIAL.
FT
    TRANSMEM
               167
                      187
                               POTENTIAL.
FT
    DOMAIN
                49
                      237
                               RETICULON.
SO
    SEQUENCE
               237 AA; 25428 MW; EB60A0A7AC45F0DE CRC64;
  Query Match
                        10.7%; Score 625.5; DB 1; Length 237;
  Best Local Similarity 59.1%; Pred. No. 1.4e-17;
  Matches 114; Conservative 41; Mismatches
                                              37: Indels
                                                            1; Gaps
                                                                       1;
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Qу
                45 SSCAVHDLIFWRDVKKTGFVFGTTLIMLLSLAAFSVISVVSYLILALLSVTISFRVYKSV 104
Db
Qу
        1032 IQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLK 1091
             Db
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        1092 FAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKI 1151
Qу
              | : ||
Db
         165 LAVFMWLMTYVGAVFNGITLLILAELLVFSVPIVYEKYKTQIDHYVGIARDQTKSIVEKI 224
        1152 QAKIPGL-KRKAD 1163
Qу
             Dh
         225 QAKLPGIAKKKAE 237
RESULT 8
RTN2 MOUSE
    RTN2 MOUSE
                  STANDARD;
                                PRT:
                                       471 AA.
    070622; 070620;
AC
DT
    16-OCT-2001 (Rel. 40, Created)
DT
    16-OCT-2001 (Rel. 40, Last sequence update)
DT
    28-FEB-2003 (Rel. 41, Last annotation update)
    Reticulon protein 2 (Neuroendocrine-specific protein-like 1) (NSP-like
DE
    protein 1) (NSPLI).
    RTN2 OR NSPL1.
GN
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
    [1]
RΡ
    SEQUENCE FROM N.A. (ISOFORMS 1 AND 2), AND TISSUE SPECIFICITY.
RC
    STRAIN=FVB/N, and 129/Sv; TISSUE=Cerebellum, and Skeletal muscle:
RX
    MEDLINE=98191726; PubMed=9530622;
RA
    Geisler J.G., Stubbs L.J., Wasserman W.W., Mucenski M.L.;
RT
    "Molecular cloning of a novel mouse gene with predominant muscle and
RT
    neural expression.";
    Mamm. Genome 9:274-282(1998).
RL
CC
    -!- SUBCELLULAR LOCATION: Membrane-bound. Endoplasmic reticulum
CC
        (Potential).
CC
    -!- ALTERNATIVE PRODUCTS:
CC
        Event=Alternative splicing; Named isoforms=2;
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CC
        Name=1; Synonyms=Brain;
CC
          IsoId=070622-1; Sequence=Displayed;
CC
        Name=2; Synonyms=Muscle;
CC
          IsoId=070622-2; Sequence=VSP 005650, VSP 005651;
    -!- TISSUE SPECIFICITY: EXPRESSED PREDOMINANTLY IN NEURAL AND MUSCULAR
CC
CC
        TISSUES.
CC
    -!- SIMILARITY: Contains 1 reticulon domain.
CC
CC
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DR
    EMBL; AF038537; AAC14906.1; -.
DR
    EMBL; AF038537; AAC14907.1; -.
    EMBL; AF038538; AAC14908.1; -.
DR
    EMBL; AF038539; AAC14909.1; -.
DR
DR
    EMBL; AF093624; AAD13195.1; -.
DR
    MGD; MGI:107612; Rtn2.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
    PROSITE; PS50845; RETICULON; 1.
KW
    Endoplasmic reticulum; Alternative splicing; Transmembrane.
FT
    TRANSMEM
               295
                    315
                             POTENTIAL.
FT
    DOMAIN
               272
                     471
                              RETICULON.
FT
    VARSPLIC
               1
                     267
                              Missing (in isoform 2).
FT
                              /FTId=VSP 005650.
FT
    VARSPLIC 268 271
                              PLLL -> MGSK (in isoform 2).
FT
                             /FTId=VSP 005651.
SO
    SEQUENCE 471 AA; 51346 MW; 9BBD8F372CF63AD3 CRC64;
 Query Match
                       8.7%; Score 509; DB 1; Length 471;
 Best Local Similarity 28.6%; Pred. No. 8.3e-13;
 Matches 146; Conservative 84; Mismatches 175; Indels 106; Gaps
QУ
         690 PDFSNYSEIAKFEKSVPEHAELVEDSSP--ESEPVDLFSDDSIPEVPQTQEEAVMLMKES 747
            Db
          6 PVFAHCKEAPSTASSTPDSTEGGNDDSDFRELHTAREFSED------ 46
         748 LTEVSETVAQH-------KEERLSASPQELGKPYLESFQPNLHST 785
QУ
                                         47 -- EEEETTSQDWGTPRELTFSYIAFDGVVGSGGRRDSVVRRPRPQGRSVSEPRDPPQQSG 104
Db
Qу
         786 KDAASNDIPTLTKK-----EKISLQMEEFNTAIYSNDDLLSSKEDK 826
              : ||:|::
                                         |:: |:::: : | | | | | |
Db
         105 LGDSLESIPSLSQSPEPGRRGDPDPVPPAERPLEELRLRLDQLGWVVRS----AGSGED- 159
Qу
         827 IKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCL 886
          Db
         160 ----SATSSSTPLE--NEEPDGLEASE----AGEETNLEL----RLAO------SL 195
         887 ELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTK 946
Qу
             Db
        196 HLQLEVLTPQLSPSSGTPQAHTPSPQRSQDSNS--GPDDEPLLNVVEEHWRLLEQEPITA 253
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Qу
         947 EAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFS 1006
                     |:::|:|
                                       Db
         254 QC----LDSTDQSEFMLEPLL------LVADLLYWKDTRTSGAVFTGLMASLLCLLHFS 302
        1007 IVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSA 1066
Qу
             303 IVSVAAHLALLGLCATISLRVYRKVLQAVHRGDGTNPFQAYLDMDLTLTREQTERLSQQI 362
Db
QУ
        1067 LGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIY 1126
               363 ASHVVSTATQLRHFFLVEDLVDSLKLALLFYILTFVGAIFNGLTLVILGVVALFTVPLLY 422
Db
Qу
        1127 ERHQVQIDHYLGLANKSVKDAMAKIQAKIPG 1157
              Db
         423 RQHQAQIDQYVGLVTNQLSHIKAKIRAKIPG 453
RESULT 9
RTN2 HUMAN
                                PRT;
    RTN2 HUMAN
                  STANDARD;
                                      545 AA.
AC
    075298; 060509;
DT
    16-OCT-2001 (Rel. 40, Created)
    16-OCT-2001 (Rel. 40, Last sequence update)
    15-SEP-2003 (Rel. 42, Last annotation update)
    Reticulon protein 2 (Neuroendocrine-specific protein-like 1) (NSP-like
DE
DE
    protein 1) (NSPLI).
GN
    RTN2 OR NSPL1.
OS
    Homo sapiens (Human).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
    NCBI TaxID=9606:
RN
    SEQUENCE FROM N.A., ALTERNATIVE SPLICING, AND TISSUE SPECIFICITY.
RΡ
RC
    TISSUE=Lung carcinoma;
    MEDLINE=98360096; PubMed=9693037;
RX
    Roebroek A.J.M., Contreras B., Pauli I.G.L., Van de Ven W.J.M.;
RA
RT
    "cDNA cloning, genomic organization, and expression of the human RTN2
    gene, a member of a gene family encoding reticulons.";
RL
    Genomics 51:98-106(1998).
RN
    [2]
RP
    SEQUENCE OF 108-545 FROM N.A. (ISOFORM RTN2-B).
RC
    TISSUE=Brain;
RX
    MEDLINE=98191726; PubMed=9530622;
RA
    Geisler J.G., Stubbs L.J., Wasserman W.W., Mucenski M.L.;
    "Molecular cloning of a novel mouse gene with predominant muscle and
RT
    neural expression.";
    Mamm. Genome 9:274-282(1998).
RL
CC
    -!- SUBCELLULAR LOCATION: Integral membrane protein. Endoplasmic
        reticulum (Potential).
CC
CC
    -!- ALTERNATIVE PRODUCTS:
CC
        Event=Alternative splicing; Named isoforms=2;
CC
        Name=RTN2-A;
CC
          IsoId=075298-1; Sequence=Displayed;
CC
          Note=Isoform RTN2-C is produced by alternative initiation at
CC
          Met-341 of isoform RTN2-A;
CC
        Name=RTN2-B;
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CC
          IsoId=075298-2; Sequence=VSP 005649;
CC
        Event=Alternative initiation;
CC
          Comment=2 isoforms, RTN2-A (shown here) and RTN2-C, are produced
CC
          by alternative initiation at Met-1 and Met-341;
CC
    -!- TISSUE SPECIFICITY: ISOFORM RTN2-C IS HIGHLY EXPRESSED IN SKELETAL
CC
        MUSCLE.
CC
    -!- SIMILARITY: Contains 1 reticulon domain.
CC
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CC
    EMBL; AF004222; AAC32542.1; -.
DR
DR
    EMBL; AF004223; AAC32543.1; -.
DR
    EMBL; AF004224; AAC32544.1; -.
DR
    EMBL; AF038540; AAC14910.1; -.
DR
    Genew; HGNC:10468; RTN2.
    MIM; 603183; -.
DR
DR
    GO; GO:0030176; C:endoplasmic reticulum membrane, intrinsic p. . .; NAS.
    GO; GO:0004871; F:signal transducer activity; NAS.
DR
    GO; GO:0007165; P:signal transduction; NAS.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
DR
    PROSITE; PS50845; RETICULON; 1.
KW
    Endoplasmic reticulum; Alternative splicing; Transmembrane;
KW
    Alternative initiation.
FT
    CHAIN
                               RETICULON PROTEIN 2, ISOFORM RTN2-A.
                     545
FT
    CHAIN
               341
                     545
                              RETICULON PROTEIN 2, ISOFORM RTN2-C.
FT
    INIT MET
               341
                    341
                               FOR ISOFORM RTN2-C.
FT
    TRANSMEM
               368
                     388
                              POTENTIAL.
FT
    TRANSMEM
               463
                     483
                              POTENTIAL.
FT
               345 545
    DOMAIN
                              RETICULON.
FT
    VARSPLIC 272 344
                              Missing (in isoform RTN2-B).
FT
                              /FTId=VSP 005649.
SO
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 Query Match
                        8.4%; Score 490; DB 1; Length 545;
 Best Local Similarity 28.3%; Pred. No. 5.2e-12;
 Matches 160; Conservative 76; Mismatches 202; Indels 128; Gaps
                                                                     17:
         668 EAPYISIACDLIKETKLSTEPSPDFSNYSEI---AKFEKSVPEHAELVEDSSPESEPVDL 724
Qу
                         Db
          13 EAP----STASSTPDSTEGGNDDSDFRELHTAREFSEEDEEETTSQDWGTPRELTFSY 66
         725 FSDDSIPEVPQTQEEAVMLMKESLTEVSETVAOHKEERLSASPQELGKPYL-ESFOPNLH 783
Qу
             67 IAFDGVVGSGGRRDSTARRPRPQGRSVSEPRDQHPQPSLGDSLESI--PSLSQSPEPGRR 124
Db
         784 STKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIID 843
Qу
               Db
         125 GDPDTAP---PSERPLEDLRLRLDHLGWVARGT----GSGED-----SSTSSSTPLE--- 169
Оv
         844 EFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIOSGADSLPCLELPCDLSFKNIYPKDEV 903
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|: | : | | | : |
                                                   | | | | | | | | |
         170 -------DEEP---QEPNRLETGEAGE------EL--DLRLRLAQPSSPE 201
Dh
         904 HVSDEFSENRSSVSKASISP----SNVSALEPOTE----- 934
Qу
                    : : || || || ||
         202 VLTPQLSPGSGTPQAGTPSPSRSRDSNSGPEEPLLEEEEKOWGPLEREPVRGOCLDSTDO 261
Db
         935 -----MGSIVK--SKSLTKEAEKKLP------ 954
Qу
                     :|: :: || :|
Db
         262 LEFTVEPRLLGTAMEWLKTSLILLAVYKTVPILELSPPLWTAIGWVORGPTPPTPVLRVLL 321
         955 DTEKEDRSLSAVLSAELSK---TSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVT 1011
QУ
               322 KWAKSPRS-SGVPSLSLGADMGSKVADLLYWKDTRTSGVVFTGLMVSLLCLLHFSIVSVA 380
Db
Qу
        1012 AYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVOKYSNSALGHVN 1071
             381 AHLALLLLCGTISLRVYRKVLQAVHRGDGANPFQAYLDVDLTLTREQTERLSHQITSRVV 440
Db
        1072 STIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQV 1131
Qу
                Db
         441 SAATQLRHFFLVEDLVDSLKLALLFYILTFVGAIFNGLTLLILGVIGLFTIPLLYRQHQA 500
QУ
        1132 QIDHYLGLANKSVKDAMAKIQAKIPG 1157
             | | | | : | | :
                           Db
         501 QIDQYVGLVTNQLSHIKAKIRAKIPG 526
RESULT 10
PCLO HUMAN
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                 STANDARD;
                              PRT; 5147 AA.
AC
    Q9Y6V0; O43373; O60305; Q9BVC8; Q9UIV2; Q9Y6U9;
    28-FEB-2003 (Rel. 41, Created)
DT
    28-FEB-2003 (Rel. 41, Last sequence update)
DT
    15-SEP-2003 (Rel. 42, Last annotation update)
DE
    Piccolo protein (Aczonin) (Fragments).
GN
    PCLO OR ACZ OR KIAA0559.
OS
    Homo sapiens (Human).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OX
    NCBI TaxID=9606;
RN
    [1]
RΡ
    SEQUENCE OF 1-759 FROM N.A.
    TISSUE=Brain;
RC
RX
    MEDLINE=99439764; PubMed=10508862;
RA
    Wang X., Kibschull M., Laue M.M., Lichte B., Petrasch-Parwez E.,
    Kilimann M.W.;
RA
RT
    "Aczonin, a 550-kd putative scaffolding protein of presynaptic active
RT
    zones, shares homology regions with rim and bassoon and binds
RT
    profilin.";
RL
    J. Cell Biol. 147:151-162(1999).
RN
    [2]
RΡ
    SEQUENCE OF 552-4404 FROM N.A.
    Kraemer J., Wollam C., Wohldmann P., McGrane B.;
RA
RL
    Submitted (DEC-1999) to the EMBL/GenBank/DDBJ databases.
RN
    [3]
RP
    SEQUENCE OF 3619-5147 FROM N.A. (ISOFORM 2).
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RC
     TISSUE=Brain;
RX
     MEDLINE=98290545; PubMed=9628581;
RA
     Nagase T., Ishikawa K.-I., Miyajima N., Tanaka A., Kotani H.,
RA
     Nomura N., Ohara O.;
RT
     "Prediction of the coding sequences of unidentified human genes. IX.
RT
     The complete sequences of 100 new cDNA clones from brain which can
RT
     code for large proteins in vitro.";
RL
     DNA Res. 5:31-39(1998).
RN
RP
     SEQUENCE OF 4405-4439 FROM N.A.
RC
     TISSUE=Placenta;
     MEDLINE=22388257; PubMed=12477932;
RX
RA
     Strausberg R.L., Feingold E.A., Grouse L.H., Derge J.G.,
RA
     Klausner R.D., Collins F.S., Wagner L., Shenmen C.M., Schuler G.D.,
RA
     Altschul S.F., Zeeberg B., Buetow K.H., Schaefer C.F., Bhat N.K.,
     Hopkins R.F., Jordan H., Moore T., Max S.I., Wang J., Hsieh F.,
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     Diatchenko L., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
RA
RA
     Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Scheetz T.E.,
RA
     Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
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     Raha S.S., Loquellano N.A., Peters G.J., Abramson R.D., Mullahy S.J.,
RA
     Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
RA
     Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
RA
     Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
     Fahey J., Helton E., Ketteman M., Madan A., Rodrigues S., Sanchez A.,
RA
     Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
RA
     Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
RA
     Rodriguez A.C., Grimwood J., Schmutz J., Myers R.M.,
RA
     Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.E.,
RA 
     Schnerch A., Schein J.E., Jones S.J.M., Marra M.A.;
RT
     "Generation and initial analysis of more than 15,000 full-length
RT
     human and mouse cDNA sequences.";
RL
     Proc. Natl. Acad. Sci. U.S.A. 99:16899-16903(2002).
RN
RΡ
     SEQUENCE OF 4405-5147 FROM N.A.
RA
     Kalicki J., Elliott G.;
RL
     Submitted (FEB-1998) to the EMBL/GenBank/DDBJ databases.
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     -!- FUNCTION: May act as a scaffolding protein involved in the
         organization of synaptic active zones and in synaptic vesicle
CC
CC
         trafficking (By similarity).
CC
     -!- SUBUNIT: Interacts with Rabacl/Pral and profilin (By similarity).
CC
     -!- SUBCELLULAR LOCATION: Concentrated at the presynaptic side of
CC
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CC
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CC
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CC
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CC
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CC
CC
         phospholipids. Calcium binds with low affinity but with high
CC
         specificity and induces a large conformational change.
CC
     -!- SIMILARITY: Contains 2 C2 domains.
CC
     -!- SIMILARITY: Contains 1 PDZ/DHR domain.
CC
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CC
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     or send an email to license@isb-sib.ch).
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     GO; GO:0045202; C:synaptic junction; ISS.
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     GO; GO:0005544; F:calcium-dependent phospholipid binding acti. . .; ISS.
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DR
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     PRINTS; PR00360; C2DOMAIN.
DR
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                   1
                          1
FT
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                        465
                                   10 X 10 AA TANDEM APPROXIMATE REPEATS OF
FТ
                                   P-A-K-P-Q-P-Q-O-P-X.
FT
     ZN FING
                 499
                        523
                                   C4-TYPE (POTENTIAL).
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                1010
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                2300
                       2325
                                   POLY-PRO.
FT
     DOMAIN
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                                   PDZ.
FT
     DOMAIN
                                   C2 DOMAIN 1.
                4544
                       4633
FT
     DOMAIN
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                       5121
                                   C2 DOMAIN 2.
FT
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                4404
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                       4404
FT
                                   QTGKLMEG (in isoform 2).
FT
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SO
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	Db	251	:	296
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	Db	297	AQPSGLTKPLAQQPGTVKPPVQPPGTTKPPAQPLGPAKPPAQQTGSEKPSSEQPGPKA	354
	Qy	102	PERQPSWERSPA-APAPSLP	138
	Db	355	LAQPPGVGKTPAQQPGPAKPPTQQVGTPKPLAQQPGLQSPAKAPGPTKTPAQTKPPSQQP	414
	Qy	139	RPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIM :	195
	Db	415	GSTKPPPQQPGPAKPSPQQPGSTKPPSQQPGSAKPSAQQPSPAKPSA	461
	Qy	196	DLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPL-STVSFKEH-GYLGNLSAVS : : : : : : : : : : : : : :	247
	Dр	462	QQFTKPVSQTGFGKPLQPPTVSPSAKQPPSQGLPKTICPLCNTTELLLHVPEKANFNTCT	521
	QУ	248	SSEGTIPERATNPFVNRDL: :	277
	Db	522	ECQTTVCSLCGFNPNPHLTEAKEWLCLNCQMKRALGGDLAPVPSSPQPKLKTAPVTTTSA	581
	Qу	278	AEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGK	337
	Db	582	VSKSSPQPQQTSPKKDAAPKQDLSKAPEPKKPPPLVKQPTLHGSPSAKAKQPPEADSLSK	641
	Qу	338	EDRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVES : : : : : : : : : : : : : : : :	397
	Db	642	PAPPKEPSVPSEQDKAPVADDKP-KQPKMVKPTTDLVSSSSATTKP	686
	Qу	398	KVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTT::::::::::::::::::::::::::::::::	454
	Db	687	DIPSSKVQSQAEEKTTPPLKTDSAKPSQSFPPTGEKVTPFDSKAIPRP	734
	Qу	455	ANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPK	490
	Db	735	ASDSKIISHPGPSSESKGQKQVDPVQKKEEPKKAQTKMSPKPDAKPMPKGSPTPPGP	791
	Qу	491	TSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEAT : : : :	544
	Db	792	RPTAGQTVPTPQQSPKPQEQSRRFSLNLGSITDAPKSQPTTPQETVT	838
,	Qy	545	GTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSA : : : : : : : : : : : : : : : :	604
	Db	839	GKLFGFGASI-FSQASNLISTAGQPGPHSQSGPGAPMKQAPAPSQPPTS	886
•	Qy	605	GASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEE : : : : : : :	639
3	Db	887	QGPPKSTGQAPPAPAKSIPVKKETKAPAAEKLEPKAEQAPTVKRTETEKKPPPIKD	942
(Qу	640	AMNVALKALGTKEGIKEPESFNAAVQE	666

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| |: | : | |
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        667 TEAPYISIAC------DLIK-----ETKLSTEPSPDFSNYSEIAKFEKSVPEHA 709
Qу
              :| : : : | |
Db
        1003 NPTPHLTENCQTQRAISGQLGDIRKMPPAPSGPKASPMPVP----TESSSQKTAVPPOV 1057
        710 ELV----EDSSPESEPV-----DLFSDDSIPEV---PQTQEEAVMLMKESLTEVSETVA 756
Qу
            Db
        1058 KLVKKQEQEVKTEAEKVILEKVKETLSMEKIPPMVTTDQKQEES-KLEKDKASALQEKKP 1116
QУ
        757 QHKEERLSASPQEL----GKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLOMEEFNTA 812
             1117 LPEEKKLIPEEEKIRSEEKKPLLEEKKPTPEDKK------LLPEAKTSAPEEQ---- 1163
Db
Qу
        813 IYSNDDLLSSK----EDKI------KESETFSDSSPIEIIDEFPTFVSAKDDS- 855
                Db
       1164 ---KHDLLKSQVQIAEEKLEGRVAPKTVQEGKQPQTKMEGLP----SGTPQSLPKEDDKT 1216
Qу
        856 -----SGADSLPCLELPCD 891
                    | || : | |||: :: |
                                         | | | :|
       1217 TKTIKEQPQPPCTAKPDQEKE-DDKSDTSSSQQPKSPQGLSDTGYSSDGISSSLGEIP-- 1273
Db
        892 LSFKNIYPKDEVHV----SDEFSENRSSVSK---ASISPSNVSALEPOTEMGSIVKSKS 943
Qу
             Dh
       1274 ----SLIPTDEKDILKGLKKDSFSQESSPSSPSDLAKLESTVLSILEAQASTLADEKSEK 1329
Qy
        944 LTK--EAEKKLPSDTEKEDRSLSAVLSAELSKTSV 976
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DT
    28-FEB-2003 (Rel. 41, Created)
DT
    28-FEB-2003 (Rel. 41, Last sequence update)
    28-FEB-2003 (Rel. 41, Last annotation update)
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OS
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    MEDLINE=99439764; PubMed=10508862;
RA
    Wang X., Kibschull M., Laue M.M., Lichte B., Petrasch-Parwez E.,
RA
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    "Aczonin, a 550-kd putative scaffolding protein of presynaptic active
RT
RT
    zones, shares homology regions with rim and bassoon and binds
RT
    profilin.";
RL
    J. Cell Biol. 147:151-162(1999).
    -!- FUNCTION: May act as a scaffolding protein involved in the
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CC
        organization of synaptic active zones and in synaptic vesicle
CC
        trafficking (By similarity).
CC
     -!- SUBUNIT: Interacts with Rabacl/Pral and profilin (By similarity).
CC
     -!- SUBCELLULAR LOCATION: Concentrated at the presynaptic side of
CC
        synaptic junctions (By similarity).
CC
     -!- DOMAIN: C2 domain 1 is involved in binding calcium and
CC
        phospholipids. Calcium binds with low affinity but with high
CC
        specificity and induces a large conformational change.
CC
     -!- SIMILARITY: Contains 2 C2 domains.
     -!- SIMILARITY: Contains 1 PDZ/DHR domain.
CC
CC
CC
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    entities requires a license agreement (See http://www.isb-sib.ch/announce/
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    or send an email to license@isb-sib.ch).
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DR
    GO; GO:0005509; F:calcium ion binding activity; ISS.
DR
    GO; GO:0005544; F:calcium-dependent phospholipid binding acti. . .; ISS.
    GO; GO:0005522; F:profilin binding activity; ISS.
    GO; GO:0007010; P:cytoskeleton organization and biogenesis; ISS.
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    GO; GO:0016080; P:synaptic vesicle targeting; ISS.
DR
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    InterPro; IPR000008; C2.
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DR
    Pfam; PF00168; C2; 2.
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    Pfam; PF00595; PDZ; 1.
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    SMART; SM00239; C2; 2.
DR
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DR
DR
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DR
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KW
KW
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FT
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FT
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FT
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               4627
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                                                                        56;
Qу
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Qy	110	RSPAAPAPSLPPAAAVLPSKLPEDDEPPAR	140
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Qу	197	LMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEET : : : : :	256
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Qy	346	KTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESK	398
Db	474	SQQADSKKPVPQKKQPSMPGSPPVKSKQTHAEPSDTGQQI-DSTPKSDQVKPTQA	527
Qy	399	VDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSA : : : : : : :	449
Db	528	EEKQNQPSIQKPTMDTVPTSAAPGVKQDLADPQSPSTQQKVTDSPMPET	576
Qy	450	TESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLV : : : : :	497
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Qy	498	A-VQDSEADYVTTDTLSKVTEAAVSNMPE	525
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Qy	526	GLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQ	564
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Qy	565	ESLYPTAQLCPSFEEAEATPSP-VLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPP	621
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Qy	622	VSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAA::	663
Db	806	LAKDSKHQTAEAKKPAELSEQEKASQPKVSCPLCKTGLNIGSKDPPNFNTCTECKKVVCN	865
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        773 -----PYLESFQPNLHSTKDA---ASNDIPTLTKKEKISLQMEEFNTAIYSN---- 816
Qу
                    1042 IPSADKILHRLQKEDPKLQQMKMAKALSADKIQPEAQKEDVQLQEVRLSKAVSADKIQHG 1101
Db
        817 --DDL------ 851
Qу
              Db
        1102 IQKDLNLQHVKIEKTSSVEKIQEAQKESKLQQDKLPKTLSEDKIPATVSSDHKKLLSKSE 1161
Qу
        852 KDDSPKLAKEYTDLEVSDKSEIANIQSG--ADSLPCLELPCDLSFKNIYPKDEVHVSDE- 908
            1162 EDKKPELLEKSTPHPKDKKEQITAETTGHITEQKVEVEAPCD----KLHEKKQEDVKKED 1217
Db
Qу
        909 ----FSENRSSVSKA--SISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRS 962
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        963 LS 964
Db
        1273 KS 1274
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AC
    Q02910;
DT
    01-OCT-1993 (Rel. 27, Created)
DT
    01-OCT-1993 (Rel. 27, Last sequence update)
DT
    15-SEP-2003 (Rel. 42, Last annotation update)
DE
    Calphotin.
GN
    CPN OR CAP.
OS
    Drosophila melanogaster (Fruit fly).
OC
    Eukaryota; Metazoa; Arthropoda; Hexapoda; Insecta; Pterygota;
OC
    Neoptera; Endopterygota; Diptera; Brachycera; Muscomorpha;
OC
    Ephydroidea; Drosophilidae; Drosophila.
OX
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RN
    [1]
RΡ
    SEQUENCE FROM N.A.
RC
    STRAIN=Canton-S;
RX
    MEDLINE=93165729; PubMed=8094559;
RA
    Martin J.H., Benzer S., Rudnicka M., Miller C.A.;
RT
    "Calphotin: a Drosophila photoreceptor cell calcium-binding protein.";
RL
    Proc. Natl. Acad. Sci. U.S.A. 90:1531-1535(1993).
RN
    [2]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=Canton-S;
    MEDLINE=93165730; PubMed=8434015;
RX
RA '
    Ballinger D.G., Xue N., Harshman K.D.;
RT
    "A Drosophila photoreceptor cell-specific protein, calphotin, binds
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681 -----ETKLSTEPSPDFSNYSEIAKFEKSVP--EHAEL------VEDSS 716

Qу

```
calcium and contains a leucine zipper.";
RL
     Proc. Natl. Acad. Sci. U.S.A. 90:1536-1540(1993).
     -!- FUNCTION: Might function as a calcium-sequestering "sponge" to
CC
CC
        regulate the amount of free cytoplasmic calcium. It binds 0.3 mole
CC
        of Ca(2+) per mole of protein.
CC
     -!- SUBUNIT: Homodimer (Probable).
CC
     -!- SUBCELLULAR LOCATION: CYTOPLASMIC; HYPODENSE COMPARTMENT.
CC
     -!- TISSUE SPECIFICITY: SOMA AND AXONS OF PHOTORECEPTOR CELLS OF
CC
        COMPOUND EYES AND OCELLI.
CC
     -!- DEVELOPMENTAL STAGE: EXPRESSED EARLY IN PHOTORECEPTOR CELL
CC
        DEVELOPMENT.
CC
CC
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     or send an email to license@isb-sib.ch).
CC
     ______
     EMBL; L02111; AAA28405.1; -.
DR
     EMBL; L05080; AAA28420.1; -.
DR
DR
     PIR; A47282; A47282.
DR
     PIR; A47283; A47283.
     FlyBase; FBgn0010218; Cpn.
DR
     GO; GO:0005509; F:calcium ion binding activity; IDA.
KW
     Calcium-binding.
FT
     CONFLICT
                36
                       36
                               A -> AVAPAVVA (IN REF. 2).
FT
     CONFLICT
                 43
                       43
                               I -> T (IN REF. 2).
FT
     CONFLICT
                64
                       64
                               I -> V (IN REF. 2).
FT
     CONFLICT
                76
                       76
                               T \rightarrow A (IN REF. 2).
FT
     CONFLICT
              100 100
                               P -> PP (IN REF. 2).
FT
     CONFLICT
               126 127
                               VQ \rightarrow AP (IN REF. 2).
              154 154
160 160
534 534
699 699
FT
    CONFLICT
                               I -> V (IN REF. 2).
FT
    CONFLICT
                               S \rightarrow T (IN REF. 2).
FT
    CONFLICT
                              A \rightarrow E (IN REF. 2).
FT
    CONFLICT
                               I -> T (IN REF. 2).
FT
    CONFLICT
               703
                     703
                               V \rightarrow L (IN REF. 2).
FT
    CONFLICT
               721
                     721
                              D -> E (IN REF. 2).
    SEQUENCE 865 AA; 84781 MW; 2110417E0B0E7CFE CRC64;
  Query Match
                         5.1%; Score 295.5; DB 1; Length 865;
  Best Local Similarity 21.3%; Pred. No. 0.00021;
 Matches 217; Conservative 127; Mismatches 379; Indels 295; Gaps
          62 AAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPER----QPSWERSPAAPAP 117
Qу
             Db
          11 SAPVAAPVTPSAVAAPVQVVSPAAVAPAPAAPIAVTPVAPPPTLASVQPATVTIP-APAP 69
Qу
         118 ----SLPPAAAVLPSKLPEDDEPPARPPPPPPAGA------SPLAEPAAPPSTP---- 161
                                                :|:| | |
                 Db
          70 IAAASVTPVASVAPPVVAAPTPPAASPVSTPVAVAQIPVAVSAPVAPPVAATPTPVVQIP 129
         162 -AAPKRRGSGSVDETLFALP--AASEP----VIPSSAEKIMDLMEQPGNT---VSSGQED 211
Ov
                   Db
         130 VAAP-----VIATPPVAASAPTPAAVTPVISPVIASPPVVPANTTVPVAAPVAA 178
```

RT

Qy	212	FPSVLLETAASL-PSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEASKELPERATN : : : : : :	270
Db	179	VPAAVPVVAPVLAPAVAPAVAPVVAETPAPPPVAEIPVATIPECVAPLIPEVSVVATK	236
Qy		PFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSP	
Db	237		263
Qy	331	QESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLA	390
Db	264	: :::: HVSVAPAVETAVVAPV	279
Qy	391	ARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASF	446
Db	280	: : : : : : :: : : : -SASTEPPVAAATLTTAPETPALAPVVAESQVAANTVVATPPTPAPEPETIAPPVVAETP	338
Qy	447	TSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADY	506
Db	339	:: :: : : : : : : : : : : : : : :	381
Qy	507	VTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQES	566
Db	382	: : :	414
Qy	567	LYP-TAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAP	619
Db	415	: : : :: : : : : LPPVAAEPVPAVVAEETPETPAPASAPVTI-AALDIPEVAPVIAAPSDAPAEAPSAAA	471
Qy		PPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQETEAPYISIACD	
Db	472		498
Qy	678	LIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFSD-DSIP	731
Db	499	:: : : : : : : : :	558
Qy	732	EVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTKDAASN	791
Db	559	: : : : : : : : : : : : : : : : : :	614
Qу	792	DIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSA	851
Db	615	:	651
Qy	852	KDDSPKLAKEYTDLEV-SDKSEIANIQSGADSLPCLELPCDLSFKNIYP-	899
Db	652	: : : :: : : KVLDPAITEAPVTTQEPDVANINDGAPATEITTPAVEIVTAAAEVSDIAIPVIDPP	707
Qу	900	-KDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMG	936
Db	708	: : : :: : : : VPQEIAVA-EIPETDTKPAEVIVEQSTIPIEAPVPEVSKYAEPVISEAPAAEVPITAGDN	766
Qy	937	SIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGV 99	0
Db	767	: : : : : : : :: PDNTSVGISEVVPTIAEKPVEEVPTSEIPEQSSSPSDSVPVAKITPLLRDLQTTDV 82	!2

```
RESULT 13
MAPB RAT
     MAPB RAT
                    STANDARD;
                                    PRT:
                                         2459 AA.
AC
     P15205; Q62958; Q9ER21; Q9QW92;
DT
     01-APR-1990 (Rel. 14, Created)
     16-OCT-2001 (Rel. 40, Last sequence update)
DT
     28-FEB-2003 (Rel. 41, Last annotation update)
DE
     Microtubule-associated protein 1B (MAP 1B) (Neuraxin) [Contains: MAP1
DE
     light chain LC1].
GN
     MAP1B.
OS
     Rattus norvegicus (Rat).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
     Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.
OX
     NCBI TaxID=10116;
RN
     [1]
RP
     SEQUENCE OF 1-142 FROM N.A.
RC
     STRAIN=Sprague-Dawley; TISSUE=Testis;
RX
     MEDLINE=96257242; PubMed=8666295;
RA
     Liu D., Fischer I.;
RT
     "Isolation and sequencing of the 5' end of the rat microtubule-
RT
     associated protein (MAP1B) -encoding cDNA.";
RL
     Gene 172:307-308(1996).
RN
     [2]
RP
     SEQUENCE OF 96-2459 FROM N.A., DOMAIN, AND INDUCTION.
RC
     STRAIN=Sprague-Dawley; TISSUE=Brain, and Glial tumor;
RX
     MEDLINE=92347374; PubMed=1639092;
     Zauner W., Kratz J., Staunton J., Feick P., Wiche G.;
RA
RT
     "Identification of two distinct microtubule binding domains on
     recombinant rat MAP 1B.";
RT
RL
     Eur. J. Cell Biol. 57:66-74(1992).
RN
RΡ
     SEQUENCE OF 1541-2459 FROM N.A., AND TISSUE SPECIFICITY.
RC
     TISSUE=Spinal cord;
RX
     MEDLINE=90059871; PubMed=2555150;
RA
     Rienitz A., Grenningloh G., Hermans-Borgmeyer I., Kirsch J.,
RA
     Littauer U.Z., Prior P., Gundelfinger E.D., Schmitt B., Betz H.;
RТ
     "Neuraxin, a novel putative structural protein of the rat central
RT
     nervous system that is immunologically related to microtubule-
RT
     associated protein 5.";
RL
     EMBO J. 8:2879-2888(1989).
RN
     [4]
RP
     DEVELOPMENTAL STAGE, AND PHOSPHORYLATION.
RX
     MEDLINE=97405699; PubMed=9260743;
RA
     Ma D., Nothias F., Boyne L.J., Fischer I.;
RT
     "Differential regulation of microtubule-associated protein 1B (MAP1B)
RT
     in rat CNS and PNS during development.";
RL
     J. Neurosci. Res. 49:319-332(1997).
CC
     -!- FUNCTION: The function of brain MAPS is essentially unknown.
CC
         Phosphorylated MAP1B may play a role in the cytoskeletal changes
CC
         that accompany neurite extension. Possibly MAP1B Binds to at least
CC
         two tubulin subunits in the polymer, and this bridging of subunits
CC
         might be involved in nucleating microtubule polymerization and in
CC
         stabilizing microtubules.
CC
     -!- SUBUNIT: 3 different light chains, LC1, LC2 and LC3, can associate
CC
         with MAP1A and MAP1B proteins.
CC
     -!- TISSUE SPECIFICITY: Nervous system (spinal cord, brain stem,
CC
         cerebellum and cerebrum). Not expressed in liver, spleen, kidney,
```

```
CC
        heart or muscle.
CC
     -!- DEVELOPMENTAL STAGE: In cerebral cortex, spinal cord and sciatic
CC
        nerve levels are high early in development but decrease during
CC
        postnatal development and are low in adults. In dorsal root
CC
        ganglia levels remain high throughout development.
CC
     -!- INDUCTION: By nerve growth factor.
CC
     -!- DOMAIN: Has a highly basic region with many copies of the sequence
CC
        KKEE and KKEI/V, repeated but not at fixed intervals, which is
CC
        responsible for the binding of MAP1B to microtubules.
CC
     -!- PTM: LCl is coexpressed with MAP1B. It is a polypeptide generated
CC
        from MAP1B by proteolytic processing. It is free to associate with
CC
        both MAP1A and MAP1B. It interacts with the amino-terminal region
CC
        of MAP1B (By similarity).
CC
     -!- PTM: Phosphorylated.
CC
     -!- SIMILARITY: TO MAPIA.
CC
     -!- CAUTION: A C-terminal fragment of this protein (residues 1597 to
CC
        2459) was originally described as neuraxin in Ref.3.
CC
     ______
CC
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     between the Swiss Institute of Bioinformatics and the EMBL outstation -
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     the European Bioinformatics Institute. There are no restrictions on its
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     use by non-profit institutions as long as its content is in no way
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     or send an email to license@isb-sib.ch).
CC
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DR
     EMBL; U52950; AAB17068.1; -.
DR
     EMBL; X60370; CAC16162.1; -.
     EMBL; X16623; CAA34620.1; ALT_SEQ.
DR
DR
     PIR; A56577; A56577.
DR
     InterPro; IPR000102; MAP1B neuraxin.
DR
     Pfam; PF00414; MAP1B neuraxin; 10.
DR
     PROSITE; PS00230; MAP1B NEURAXIN; 8.
KW
     Microtubules; Repeat; Phosphorylation.
FT
    CHAIN
                  ?
                      2459
                                MAP1 LIGHT CHAIN LC1.
FT
    REPEAT
               1869
                                MAP1B 1.
                      1885
FΤ
    REPEAT
               1886
                      1902
                                MAP1B 2.
FT
               1903
     REPEAT
                      1919
                                MAP1B 3.
FT
     REPEAT
               1920
                     1936
                                MAP1B 4.
FT
    REPEAT
               1937
                      1953
                                MAP1B 5.
FT
     REPEAT
               1954
                      1970
                                MAP1B 6.
FT
    REPEAT
               1988
                      2004
                                MAP1B 7.
FT
    REPEAT
               2005
                     2021
                                MAP1B 8.
FT
    REPEAT
               2022 2038
                                MAP1B 9.
FT
    REPEAT
               2039
                      2055
                                MAP1B 10.
FT
    DOMAIN
                559
                      1035
                                GLU-RICH.
FT
    DOMAIN
                588
                      786
                                LYS-RICH (HIGHLY BASIC, CONTAINS MANY
FT
                                KKEE AND KKEI/V REPEATS).
FT
    DOMAIN
               2224
                     2312
                                LYS-RICH.
FΤ
    CONFLICT
                127
                      127
                                M \rightarrow V (IN REF. 1).
FT
    CONFLICT
               140
                      140
                                T \rightarrow S (IN REF. 1).
FT
    CONFLICT
               2112
                      2112
                                R \rightarrow K (IN REF. 3).
FT
    CONFLICT
               2169
                      2169
                                L \rightarrow I (IN REF. 3).
SO
    SEQUENCE
               2459 AA; 269497 MW; 2E3F6872DEDB8BA2 CRC64;
 Query Match
                         5.0%; Score 291.5; DB 1; Length 2459;
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Best Local Similarity 20.0%; Pred. No. 0.0011;

Matche	es 22	0; Conservative 167; Mismatches 456; Indels 255; Gaps	44;
Qy	30	TEPEDEEDEEEEEDEEEDDEDLEELEVLERKPAAGLSAAAVPPAAAAPLLDFSSDSVPPA	89
Db	1008	:	1058
Qу	90	PRGPLPAAPPAAPERQPSWERSPAAPAPSLPPAAAVLPSKLPEDDEPPARPPPPPPAGAS	149
Db	1059	: : : :	1094
Qу	150	PLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASEPVIPSSAEKIMDLMEQPGNTVS	206
Db	1095	: :	1147
Qу	207	SGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIE	254
Db	1148	: : : : : : PSQEFVNITKYESSLYSQEYSKPVVASFNGLSDGSKTDATDGRDYNASASTISPPSSMEE	1207
Qy	255	ETLNEAS	289
Db	1208	DKFSKSALRDAYRPEETDVKTGAELDIKDVSDERLSPAKSPSLSPSPPSPIEKTPLGERS	1267
Qу	290	SSFKGSPKGESAILVENTKEEVIVRSKDKEDLVCSAALHSP	330
Db	1268	VNFSLTPNEI KASAEGEATAVVSPGVTQAVVEEHCASPEEKTLEVVSPSQSVTGSAGHTP	1327
Qу	331	-QESPVGKEDRVVSPEKTMDIFNEMQMSVVAPVREEYADFKPFEQAWEVKD	380
Db	1328	: :: : :: YYQSPTDEKSSHLPTEVTENAQAVPVSFEFTEAKDENERSSISPMDEPVPD	1378
QУ	381	TYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDASFPSTPEPVK : : : : :	434
Db	1379	SESPIEKVLSPLRSPPLIGSESAYEDFLSADDKALGRRSESPFEGKNGKQGFSDKESPVS	1438
QУ	435	DSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKI	475
Db	1439	DLTSDLYQDKQEEKRAGFIPIKEDFSPEKKASDAEIMSSQSALALDERKLGGDGS	1493
Qу	476	EERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAV	520
Db	1494	PTQVDVSQFGSFKEDTKMSISEGTVSDKSATPVDEGAEDTYSHMEGVASVSTASVAT	1550
Qу	521	SNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQES-LYPTAQLCPS	576
Db	1551	SSFPEPTTDD-VSPSLHAEVGSPHSTEVDDSLSVSVVQTPTTFQETEMSPSKEECPRPMS	1609
Qy	577	FEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQ (: : : : : : : : : :	610
Db	1610	ISPPDFSPKTAKSRTPVQDHRSEQSSMSIEFGQESPEHSLAMDFSRQSPDHPTVGAGMLH	1669
Qy	611	PSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEPESFN (661
Db	1670	ITENGPTEVDYSPSDIQDSSLSHKIPPTEEPSYTQDNDLS-ELISVSQVEASPSTSS	1725
Qy	662	AAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEP	721
Db	1726	AHTPSQIASPLQEDTLSDVVPPRDMSLYASLASEKVQSLEGEKLSPKSDI	1775

```
722 VDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYL-ESFQP 780
Qу
                  : | |
                        ::
                                 1776 SPLTPRESSPTYSPGFSDSTSGAKES-TAAYQTSSSPPIDAAAAEPYGFRSSMLFDTMQH 1834
Dh
         781 NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE 840
Qу
             1835 HLALSRDLTTSSV----EKDNGGKTPGDFNYAYQKPESTTESPDEEDYDYE--SHEKTIQ 1888
Db
         841 IIDEFPTFVSAKDDSPK----LAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKN 896
Qу
               Db
        1889 AHDVGGYYYEKTERTIKSPCDSGYSYETIEKTTKTP----EDGGYS------CEITEKT 1937
         897 IYPKDEVHVSDEFSENRSSVSKAS-----ISPSNVSALEPQTEMGSIVKSKSL 944
Qу
                1938 TRTPEEGGYSYEISEKTTRTPEVSGYTYEKTERSRRLLDDISNGYDDTEDGGHTLGDCSY 1997
Db
         945 TKEAEKKLPSDTEKEDRS 962
Oy
            : | : | : | | |
Db
        1998 SYETTEKITSFPESESYS 2015
RESULT 14
MAPB HUMAN
    MAPB HUMAN
                  STANDARD;
                             PRT; 2468 AA.
AC
    P46821;
    01-NOV-1995 (Rel. 32, Created)
DT
     01-NOV-1995 (Rel. 32, Last sequence update)
DT
    16-OCT-2001 (Rel. 40, Last annotation update)
DT
DE
    Microtubule-associated protein 1B (MAP 1B) [Contains: MAP1 light chain
DE
    LC1].
GN
    MAP1B.
    Homo sapiens (Human).
OS
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
OC
OX
    NCBI TaxID=9606;
RN
    [1]
    SEQUENCE FROM N.A.
RΡ
RC
    TISSUE=Fetal brain:
    MEDLINE=95104835; PubMed=7806212;
RΑ
    Lien L.L., Feener C., Fischbach N., Kunkel L.M.;
    "Cloning of human microtubule-associated protein 1B and the
RT
    identification of a related gene on chromosome 15.";
RT
RL
    Genomics 22:273-280(1994).
CC
    -!- FUNCTION: THE FUNCTION OF BRAIN MAPS IS ESSENTIALLY UNKNOWN.
CC
        PHOSPHORYLATED MAP1B MAY PLAY A ROLE IN THE CYTOSKELETAL CHANGES
CC
        THAT ACCOMPANY NEURITE EXTENSION. POSSIBLY MAP1B BINDS TO AT LEAST
CC
        TWO TUBULIN SUBUNITS IN THE POLYMER, AND THIS BRIDGING OF SUBUNITS
CC
        MIGHT BE INVOLVED IN NUCLEATING MICROTUBULE POLYMERIZATION AND IN
CC
        STABILIZING MICROTUBULES.
CC
    -!- SUBUNIT: 3 DIFFERENT LIGHT CHAINS, LC1, LC2 AND LC3, CAN ASSOCIATE
CC
        WITH MAP1A AND MAP1B PROTEINS.
CC
    -!- DOMAIN: Has a highly basic region with many copies of the sequence
CC
        KKEE and KKEI/V, repeated but not at fixed intervals, which is
CC
        responsible for the binding of MAP1B to microtubules.
CC
    -!- PTM: LCl is coexpressed with MAP1B. It is a polypeptide generated
```

from MAP1B by proteolytic processing. It is free to associate with

CC

```
CC
        both MAP1A and MAP1B. It interacts with the amino-terminal region
CC
        of MAP1B (By similarity).
    -!- SIMILARITY: TO MAP1A.
CC
CC
CC
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    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    entities requires a license agreement (See http://www.isb-sib.ch/announce/
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    or send an email to license@isb-sib.ch).
CC
    EMBL; L06237; AAA18904.1; -.
DR
DR
    Genew; HGNC: 6836; MAP1B.
DR
    MIM; 157129; -.
    GO; GO:0005875; C:microtubule associated complex; TAS.
DR
    InterPro; IPR000102; MAP1B_neuraxin.
DR
    Pfam; PF00414; MAP1B neuraxin; 10.
DR
    PROSITE; PS00230; MAP1B NEURAXIN; 6.
KW
    Microtubules; Repeat; Phosphorylation.
FT
    CHAIN
                   2468
                            MAP1 LIGHT CHAIN LC1.
    REPEAT
FT
              1878 1894
                            MAP1B 1.
FΤ
    REPEAT
            1895 1911
                            MAP1B 2.
    REPEAT 1912 1928
REPEAT 1929 1945
REPEAT 1946 1962
REPEAT 1963 1979
REPEAT 1997 2013
REPEAT 2014 2030
FT
                            MAP1B 3.
FT
                            MAP1B 4.
                            MAP1B 5.
FT
FT
                            MAP1B 6.
FT
                            MAP1B 7.
FT
                            MAP1B 8.
FT
    REPEAT
            2031 2047
                            MAP1B 9.
FT
    REPEAT
            2048 2064
                            MAP1B 10.
FT
    DOMAIN
             589 790
                             LYS-RICH (HIGHLY BASIC, CONTAINS MANY
FT
                             KKEE AND KKEI/V REPEATS).
    SEQUENCE 2468 AA; 270618 MW; 540839CBDF09D461 CRC64;
SQ
 Query Match
                       4.9%; Score 288; DB 1; Length 2468;
 Best Local Similarity 20.2%; Pred. No. 0.0015;
 Matches 251; Conservative 166; Mismatches 441; Indels 386; Gaps 56;
Qу
         13 STDSPPR--PPPAFKYQFVTEPEDEEDEEE-----EED-----EEEDDED 50
            Dh
         625 ATDVKPKAAKEKTVKKETKVKPEDKKEEKEKPKKEVAKKEDKTPIKKEEKPKKEEVKKEV 684
         51 LEELEVLERKPAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPERQPSWER 110
Qу
             :|:: |:| : : : : :: |:
        685 KKEIKKEEKK-----EPKKEVKKETPPKEVKKEVKKEEKKEVKKE--EK 726
Dh
Qу
        111 SPAAPAPSLPPAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRG---168
             Db
        727 EPKKEIKKLPKDAKKSSTPLSEAKKPAALKPKVPKKEESVKKDSVA----AGKPKEKGKI 782
        169 -----SGSVDETLFAL------PA----PA----ASEPVIPSSAEKIMDLME 199
Qу
                | | : |
                                          783 KVIKKEGKAAEAVAAAVGTGATTAAVMAAAGIAAIGPAKELEAERSLMSSPEDLTKDFEE 842
Dh
        200 QPGNTVSSGQEDFPSV-LLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLN 258
Qу
```

	Db	843	LKAEEVDVTKDIKPQLELIEDEEKLKETEPVEAYVIQKEREVTKGPAESPDEG-ITTTEG	901
	Qy	259	EASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTK-EEVIVRSKD	317
	Db	902		957
	Qу	318	KEDLVC-SAALHSPQESPVGKEDRVVSPEKTMDIFNEMQMSV	358
	Db	958	: : : GEEHVCVSASKHSPTEDEESAKAEADAYIREKRESVASGDDRAEEDMDEAIEKG	1011
	Qy	359	VAPVREEYADFKPFEQAWEVK-DTYEGSRDVLAARANVESKVDRKCLEDSLEQ	410
	Db	1012	EAEQSEEEADEEDKAEDAREEEYEPEKMEAEDYVMAVVDKAAEAGGAEEQYGFLTT	1067
	Qу	411	KSLGKDSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTT	454
	Db	1068	PTKQLGAQSPGREPASSIHDETLPGGSESEATASDEENREDQPEEFTATSGYTQSTIEIS	1127
	Qу	455	ANTFPLLEDHTSENKTDEKKIEERKAQIITEKTS-PKTSNPFLVAV : :	499
	Db	1128	SEPTPMDEMSTPRDVMSDETNNEETESPSQEFVNITKYESSLYSQEYSKPADVTPLNGFS	1187
	Qy	,500	QDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDL: :	556
	Db	1188	EGSKTDATDGKDYNASASTISPPSSMEEDKFSRSALRDAYCSEVKASTTLDIKDS	1242
	Qy	557	VQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIV : : : :: :::	592
	Db	1243	ISAVSSEKVSPSKSPSLSPSPPSPLEKTPLGERSVNFSLTPNEIKVSAEAEVAPVSPEVT	1302
	Qy	593	MEAPLNSLLPSAGASVVQ	610
	Db	1303	QEVVEEHCASPEDKTLEVVSPSQSVTGSAGHTPYYQSPTDEKSSHLPTEVIEKPPAVPVS	1362
	Qy	611	PSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALG	649
	Db	1363	FEFSDAKDENERASVSPMDEPVPDSESPIEKVLSPLRSPPLIGSESAYESFLSADDKASG	1422
•	Qу		TKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPD-FSNYSEIAKF	
	Db	1423	RGAESPFEEKSGKQGSPDQVSPVSEMTSTSLYQDKQ	1458
	Qу	702	EKSVPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAV	741
	Db	1459	EGKSTDFAPIKEDFGQEKKTDDVEAMSSQPALALDERKLGDVSPTQIDVSQFGSFKEDTK	1518
	Qу		MLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHSTK	
	Db	1519	MSISEGTVSDKSATPVDEGVAEDTYSHMEGVASVSTASVATSSFPEPTTDDVSPSLH	1575
	Qy	787	DAASNDIPTLTK-KEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEF : : : : :	845
	Db	1576	AEVGSPHSTEVDDSLSVSVVQTPTT-FQETEMSPSKEECPR	1615
	Qy	846	PTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKN	896
	Db	1616	PMSISPPDFSPKTAKSRTPVQDHRSEQSSM-SIEFGQES-PEQSLAMDFSRQSPDHPTVG	1673

```
Qу
         897 -----IYPKDE-VHVSDEFSENRSSVSKASISPSNV 926
                                         | | : | : |
                                                           |||:
Db
        1674 AGVLHITENGPTEVDYSPSDMQDSSLSHKIPPMEEPSYTQDNDLSELISVSQVEASPSTS 1733
Qу
         927 SALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRSLSAVLSAE 970
                 Db
         1734 SAHTP-SQIASPLQEDTLSDVAPPR-----DMSLYASLTSE 1768
RESULT 15
MAPB MOUSE
ID
    MAPB MOUSE
                   STANDARD:
                                 PRT; 2464 AA.
AC
     P14873:
     01-APR-1990 (Rel. 14, Created)
DT
DT
     01-APR-1990 (Rel. 14, Last sequence update)
     28-FEB-2003 (Rel. 41, Last annotation update)
DE
    Microtubule-associated protein 1B (MAP 1B) (MAP1.2) (MAP1(X))
DE
     [Contains: MAP1 light chain LC1].
GN
    MAP1B OR MTAP1B OR MTAP5.
OS
    Mus musculus (Mouse).
OC
     Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
     [1]
RΡ
     SEQUENCE FROM N.A., AND DOMAIN.
RC
     STRAIN=Swiss Webster; TISSUE=Brain;
    MEDLINE=90094539; PubMed=2480963;
RX
RA
    Noble M., Lewis S.A., Cowan N.J.;
     "The microtubule binding domain of microtubule-associated protein
RT
RT
    MAP1B contains a repeated sequence motif unrelated to that of MAP2
RT
    and tau.";
RL
    J. Cell Biol. 109:3367-3376(1989).
     -!- FUNCTION: THE FUNCTION OF BRAIN MAPS IS ESSENTIALLY UNKNOWN.
CC
CC
        PHOSPHORYLATED MAP1B MAY PLAY A ROLE IN THE CYTOSKELETAL CHANGES
CC
        THAT ACCOMPANY NEURITE EXTENSION. POSSIBLY MAP1B BINDS TO AT LEAST
CC
        TWO TUBULIN SUBUNITS IN THE POLYMER, AND THIS BRIDGING OF SUBUNITS
CC
        MIGHT BE INVOLVED IN NUCLEATING MICROTUBULE POLYMERIZATION AND IN
CC
        STABILIZING MICROTUBULES.
CC
     -!- SUBUNIT: 3 DIFFERENT LIGHT CHAINS, LC1, LC2 AND LC3, CAN ASSOCIATE
CC
        WITH MAPIA AND MAPIB PROTEINS.
CC
     -!- DOMAIN: Has a highly basic region with many copies of the sequence
CC
        KKEE and KKEI/V, repeated but not at fixed intervals, which is
CC
        responsible for the binding of MAP1B to microtubules.
     -!- PTM: LC1 IS COEXPRESSED WITH MAP1B. IT IS A POLYPEPTIDE GENERATED
CC
CC
        FROM MAP1B BY PROTEOLYTIC PROCESSING. IT IS FREE TO ASSOCIATE WITH
CC
        BOTH MAPIA AND MAPIB. IT INTERACTS WITH THE AMINO-TERMINAL REGION
CC
        OF MAP1B.
     -!- SIMILARITY: TO MAP1A.
CC
CC
    -----
CC
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    between the Swiss Institute of Bioinformatics and the EMBL outstation -
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    the European Bioinformatics Institute. There are no restrictions on its
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    use by non-profit institutions as long as its content is in no way
    modified and this statement is not removed. Usage by and for commercial
CC
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or send an email to license@isb-sib.ch).

CC

```
CC
DR
    EMBL; X51396; CAA35761.1; -.
    PIR; S07549; QRMSP1.
DR
DR
    MGD; MGI:1306778; Mtap1b.
DR
    GO; GO:0016358; P:dendrite morphogenesis; IMP.
    GO; GO:0001578; P:microtubule bundling; IMP.
DR
    InterPro; IPR000102; MAP1B neuraxin.
DR
DR
    Pfam; PF00414; MAP1B neuraxin; 10.
DR
    PROSITE; PS00230; MAP1B NEURAXIN; 7.
    Microtubules; Repeat; Phosphorylation.
FT
    CHAIN
                ?
                    2464
                             MAP1 LIGHT CHAIN LC1.
FT
    REPEAT
              1874
                    1890
                             MAP1B 1.
    REPEAT
FT
            1891
                   1907
                             MAP1B 2.
    REPEAT 1908 1924
REPEAT 1925 1941
REPEAT 1942 1958
REPEAT 1959 1975
    REPEAT
FT
                            MAP1B 3.
FT
                            MAP1B 4.
FT
                            MAP1B 5.
FT
                            MAP1B 6.
    REPEAT
FΤ
            1993 2009
                            MAP1B 7.
    REPEAT 2010 2026
FT
                            MAP1B 8.
              2027 2043
    REPEAT
FT
                             MAP1B 9.
    REPEAT 2044 2060
DOMAIN 589 787
FT
                            MAP1B 10.
FT
                            LYS-RICH (HIGHLY BASIC, CONTAINS MANY
FΤ
                             KKEE AND KKEI/V REPEATS).
SO
    SEQUENCE 2464 AA; 270408 MW; FBD3DD99CFDBDA87 CRC64;
 Query Match
 Query Match 4.9%; Score 284; DB 1; Length 2464; Best Local Similarity 20.7%; Pred. No. 0.0021;
 Matches 233; Conservative 159; Mismatches 424; Indels 312; Gaps 51;
Qу
         31 EPEDEEDEEEEDEEED-----DEDLEELE-----VLERKPAAG------ISAAAVP 71
            Db
        1009 EAEQSEEEGEEDKAEDAREEGYEPDKTEAEDYVMAVADKAAEAGVTEEOYGYLGTSAKO 1068
         72 PAAAAPLLDFSS----DSVPPAPRGPLPAAPPAAPEROP-----SWERSPAAPA 116
Qу
            1069 PGIQSPSREPASSIHDETLPGGSESEATASDEENREDQPEEFTATSGYTQSTIEISSEPT 1128
Db
        117 P---SLPPAAAVLPSKLPEDDEPPARP------PPPPPAGASPLAEPA-- 155
QУ
            1129 PMDEMSTPRDVMSDETNNEETESPSQEFVNITKYESSLYSQEYSKPAVASFNGLSEGSKT 1188
Db
        156 -----APPSTPAAPKRRGSGSVDETLFALPA-----ASEPVIPSSAEKIMDLMEQ 200
Оy
                   Db
        1189 DATDGKDYNASASTISPP----SSMEEDKFSKSALRDAYCSEEKELKASAE--LDIKDV 1241
        201 PGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTIEETLNEA 260
Qу
               1242 SDERLS-----PAKSPSLSP------SPPSPIEKT---- 1265
Db
        261 SKELPERATNPFVNRDLAEFS----ELEYSEMGSSFKGSPKGESAILVEN--TKEEVIVR 314
QУ
              1266 --PLGERSVN-----FSLTPNEIKVSAEGEARSVSPGVTQAVVEEHCASPEEKTLE 1314
Db
        315 SKDKEDLVCSAALHSP-QESPVGKEDRVVSPEKTMDIFNEMQMSVVA-PVREEYADFK-- 370
ΟV
       | :| |:| :|| : ||: : |: : | || |::: |
1315 VVSPSQSVTGSAGHTPYYQSP-----TDEKSSHLPTEVSENAQAVPVSFEFSEAKDE 1366
Db
```

QУ	371	PFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDS ::	417
Db	1367	:: : : : : :	1424
Qу	418	EGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKT	470
Db	1425	: : : : : : EGKNGKQGFPDRESPVSDLTSTGLYQDKQEEKSTGFIPIKEDFGPEKKTSDVETMS	1480
Qy	471	DEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEAD	505
Db	1481	: :	1538
Qy	506	YVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAI	563
Db	1539	: :: : : : : :: : : :	1597
Qy	564	QES-LYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLL	601
Db	1598	QETEMSPSKEECPRPMSISPPDFSPKTAKSRTPVQDHRSEQSSMSIEFGQESPEHSFAMD	1657
Qу	602	PSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVA	644
Db	1658	FSRQSPDHPTLGASVLHITENGPTEVDYSPCDIQDSSLSHKIPPTEEPSYTQDNDLS	1714
Qу	645	LKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKS	704
Db	1715	: : : : : : : : : : - ELISVSQVEASPSTSSAHTPSQIASPLQEDTLSDVVPPREMSLYASLA	1762
Qу	705	VPEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKE : : :	760
Db	1763	SEKVQSLEGEKLSPKSDISPLTPRESSPLYSPGFSDSTSAAKETAAAH	1810
Qу	761	ERLSASPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFN	810
Db	1811	: : : : : : : : : : : : :	1865
Qу	811	TAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLE	866
Db	1866	:: ::: : : : : : YAYQKPENAAGSPDEEDYDYESQEKTIRTHDVVRYYYEKTERTIKSPCDSGYSYETIE	1923
Qу	867	VSDKSEIANIQSGADSLPCLELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKAS	920
Db	1924	: : : : : KTTKTPEDGGYTCEITEKTTRTPEEGGYSYEISEKTTRTPEVSGYTYEK	1972
Qy	921	ISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKEDRS 962 : : : : : :	
Db	1973	TERSRRLLDDISNGYDDTEDGGHTLGDCSYSYETTEKITSFPESESYS 2020	

Search completed: December 19, 2003, 15:35:43 Job time : 23 secs

GenCore version 5.1.6 Copyright (c) 1993 - 2003 Compugen Ltd.

OM protein - protein search, using sw model

Run on:

December 19, 2003, 15:32:53; Search time 46 Seconds

(without alignments)

6524.243 Million cell updates/sec

Title:

US-09-830-972-2

Perfect score: 5848

Sequence:

1 MEDIDQSSLVSSSTDSPPRP......VKDAMAKIQAKIPGLKRKAD 1163

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched:

830525 seqs, 258052604 residues

Total number of hits satisfying chosen parameters:

830525

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database :

SPTREMBL 23:*

- 1: sp archea:*
- 2: sp_bacteria:*
- 3: sp_fungi:*
- 4: sp_human:*
- 5: sp invertebrate:*
- 6: sp mammal:*
- 7: sp mhc:*
- 8: sp_organelle:*
- 9: sp_phage:*
- 10: sp plant:*
- 11: sp_rodent:*
- 12: sp_virus:*
- 13: sp_vertebrate:*
- 14: sp unclassified:*
- 15: sp rvirus:*
- 16: sp bacteriap:*
- 17: sp archeap:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

왕 Result Query

Score Match Length DB ID Description

1	5312.5	90.8	1162	11	Q8BGM9	Q8bgm9 mus musculu
2	5307	90.7	1163	11	Q8K3G8	Q8k3g8 mus musculu
3	4501.5	77.0	1046	11	Q8BGK7	Q8bgk7 mus musculu
4	3627.5	62.0	986	4	Q8IUA4	Q8iua4 homo sapien
5	2926	50.0	639	11	Q8K290	Q8k290 mus musculu
6	1416	24.2	375	11	Q8BHF5	Q8bhf5 mus musculu
7	1314.5	22.5	356	11	Q8BH78	O8bh78 mus musculu
8	1304	22.3	357	11	Q8K3G7	Q8k3g7 mus musculu
9	1283.5	21.9	392	4	Q96B16	Q96b16 homo sapien
10	844	14.4	179	6	Q9GM33	Q9gm33 macaca fasc
11	792	13.5	780	11		Q8k4s4 mus musculu
12	788	13.5	780	11	Q8K0T0	Q8k0t0 mus musculu
13	762.5	13.0	760	13	Q90638	Q90638 gallus gall
14	700	12.0	643	11	Q8CCU2	Q8ccu2 mus musculu
15	685	11.7	199	4	Q9BQ59	Q9bq59 homo sapien
16	671	11.5	267	11	Q63765	Q63765 rattus sp.
17	669	11.4	208	13	Q90637	Q90637 gallus gall
18	625.5	10.7	236	11	Q8 VBU0	Q8vbu0 rattus norv
19	625.5	10.7	237	11	Q8 VB00	Q8c6d5 mus musculu
20	586	10.7	595	5	O9VMV9	Q9vmv9 drosophila
21	532	9.1	224	5	Q9VMW1	Q9vmv1 drosophila
22	520	8.9	202	5	Q9VMW2	Q9vmw2 drosophila
23	520	8.9	222	5	Q9VMW4	Q9vmw4 drosophila
24	520	8.9	234	5	Q9VMW3	Q9vmw3 drosophila
25	517	8.8	2484	5	Q9U347	Q9u347 caenorhabdi
26	503.5	8.6	2607	5		Q23187 caenorhabdi
27	377	6.4	2768	5	Q23187 Q9VC00	
28	347.5	5.9		5 5		Q9vc00 drosophila
29			10578		Q8ISF5	Q8isf5 caenorhabdi
30	343.5 343.5	5.9 5.9	18519	5 5	Q8ISF6 Q8ISF7	Q8isf6 caenorhabdi
		5.7	18534			To the state of th
31 32	335 331	5.7	5412	5 5	Q9W596	Q9w596 drosophila
			4900		Q9N541	Q9n541 caenorhabdi
33	328.5	5.6 5.5	5327	5	076891	076891 drosophila
34	322		7962	4	Q10465	Q10465 homo sapien
35	320	5.5	222	5	Q23188	Q23188 caenorhabdi
36	317.5	5.4 5.4	17352	5	Q95YM2	Q95ym2 procambarus
37	313.5	٠. ١	1444	5	Q9VTN2	Q9vtn2 drosophila
38	313.5	5.4	1514	5	Q8SY55	Q8sy55 drosophila
39	309.5	5.3	1852	3	Q9C2H4	Q9c2h4 neurospora
40	308	5.3	2362	5	Q9VYD1	Q9vyd1 drosophila
41	307	5:2	34350	4	Q8WZ42	Q8wz42 homo sapien
42	306	5.2	842	5	Q9VGC9	Q9vgc9 drosophila
43	306	5.2	864	5	Q9VGC8	Q9vgc8 drosophila
44	305.5	5.2	9196	5	Q8IQ87	Q8iq87 drosophila
45	304	5.2	864	5	Q95U45	Q95u45 drosophila

ALIGNMENTS

RESULT 1 Q8BGM9 ID Q8BGM9 PRELIMINARY; PRT; 1162 AA. AC Q8BGM9; DT 01-MAR-2003 (TrEMBLrel. 23, Created) DT 01-MAR-2003 (TrEMBLrel. 23, Last sequence update) DT 01-MAR-2003 (TrEMBLrel. 23, Last annotation update)

```
DE
    RTN4.
GN
    RTN4.
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
റ്
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
    [1]
RΡ
    SEQUENCE FROM N.A.
RC
    STRAIN=129/SvcJ7, and 129SvcJ7;
    Oertle T., van der Putten H., Schwab M.E.;
RA
RT
    "Genomic Structure and Functional Characterization of the Promoter
RT
    Structures of Human and Mouse Nogo/Rtn-4.";
RL
    Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
RN
    [2]
    SEQUENCE FROM N.A.
RΡ
    STRAIN=129/SvcJ7, and 129SvcJ7;
RC
RΑ
    Oertle T., Schwab M.E.;
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
RN
RP
    SEQUENCE FROM N.A.
ŔĊ
    STRAIN=129/SvcJ7;
    Van der Putten H.;
RA
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
RN
    SEQUENCE FROM N.A.
RP
    STRAIN=129SvcJ7;
RC
    Van der Putten H., Mir A.;
RA
RL
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
DR
    EMBL; AY102284; AAM73506.1; -.
DR
    EMBL; AY102286; AAM73511.1; -.
              1162 AA; 126613 MW; 855697FBEE11781F CRC64;
SO
    SEOUENCE
  Query Match
                       90.8%; Score 5312.5; DB 11; Length 1162;
  Best Local Similarity
                       91.0%; Pred. No. 3.1e-262;
 Matches 1064; Conservative 37; Mismatches
                                          55; Indels
                                                        13; Gaps
                                                                    6;
Qу
          1 MEDIDQSSLVSSSTDSPPRPPPAFKYQFVTEPEDEEDEEEEEDEEEDDEDLEELEVLERK 60
            Db
          1 MEDIDQSSLVSSSADSPPRPPPAFKYQFVTEPEDEEDEEDEE-EEEDDEDLEELEVLERK 59
         61 PAAGLSAAAVPPAAAAPLLDFSSDSVPPAPRGPLPAAPPAAPEROPSWERSPAAPAPSLP 120
Qу
            Db
         60 PAAGLSAAPVPP-AAAPLLDFSSDSVPPAPRGPLPAAPPTAPERQPSWERSPAASAPSLP 118
QУ
        121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
            Db
        119 PAAAVLPSKLPEDDEPPAR--PPAPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 176
        181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240
Qу
            Dh
        177 AASEPVIPSSAEKIMDLKEQPGNTVSSGQEDFPSVLFETAASLPSLSPLSTVSFKEHGYL 236
Qу
        241 GNLSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGES 300
            Db
        237 GNLSAVASTEGTIEETLNEASRELPERATNPFVNRESAEFSVLEYSEMGSSFNGSPKGES 296
Qу
        301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESP-----VGKEDRVVSPEKTMDIFNEMO 355
```

		1:1111111111111111111111111111111111111	
Db	297	AMLVENTKEEVIVRSKDKEDLVCSAALHNPQESPATLTKVVKEDGVMSPEKTMDIFNEMK	356
Qy	356	MSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGK	415
Db	357	MSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANMESKVDKKCFEDSLEQKGHGK	416
Qy	416	DSEGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKI	475
Db.	417	DSESRNENASFPRTPELVKDGSRAYITCDSFSSATESTAANIFPVLEDHTSENKTDEKKI	476
Qy	476	EERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEA	535
Db	477	EERKAQIITEKTSPKTSNPFLVAIHDSEADYVTTDNLSKVTEAVVATMPEGLTPDLVQEA	536
Qy	536	CESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEA	595
Db	537	CESELNEATGTKIAYETKVDLVQTSEAIQESIYPTAQLCPSFEEAEATPSPVLPDIVMEA	596
Qу	596	PLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIK	655
Db	597	PLNSLLPSTGASVAQPSASPLEVPSPVSYDGIKLEPENPPPYEEAMSVALKTSDSKEEIK	656
Qу	656	EPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDS	715
Db	657	EPESFNAAAQEAEAPYISIACDLIKETKLSTEPSPEFSNYSEIAKFEKSVPDHCELVDDS	716
Qy	716	SPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHK-EERLSASPQELGKPY	774
Db	717	SPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVTQHKHKERLSASPQEVGKPY	776
Qу	. 775	LESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFS	834
Db	777	LESFQPNLHITKDAASNEIPTLTKKETISLQMEEFNTAIYSNDDLLSSKEDKMKESETFS	836
Qy	835	DSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSF	894
Db	837	DSSPIEIIDEFPTFVSAKDDSPKEYTDLEVSNKSEIANVQSGANSLPCSELPCDLSF	893
Qy	895	KNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPS	954
Db	894	KNTYPKDEAHVSDEFSKSRSSVSKVPLLLPNVSALESQIEMGNIVKPKVLTKEAEEKLPS	953
Qу	955	DTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYI	1014
Db	954	DTEKEDRSLTAVLSAELNKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYI	1013
Qу		ALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTI	
Db		ALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTI	
Qу		KELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQID	
Db		KELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQAQID	1133
Qу	1135	HYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163	

```
RESULT 2
Q8K3G8
              PRELIMINARY;
ID
    Q8K3G8
                              PRT; 1163 AA.
AC
    Q8K3G8;
DT
    01-OCT-2002 (TrEMBLrel. 22, Created)
DT
    01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DE
    Nogo-A.
GN
    RTN4.
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
0C
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
    [1]
RΡ
    SEQUENCE FROM N.A.
RC.
    STRAIN=BALB/c:
    Jin W., Long M., Li R., Ju G.;
RA
RT
    "Cloning and expression of the mouse Nogo-A protein.";
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
DR
    EMBL; AY114152; AAM77068.1; -.
DR
    MGD; MGI:1915835; Rtn4.
DR
    InterPro; IPR003388; Reticulon.
    Pfam; PF02453; Reticulon; 1.
    PROSITE; PS50845; RETICULON; 1.
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            119 PAAAVLPSKLPEDDEPPAR--PPAPAGASPLAEPAAPPSTPAAPKRGSGSVDETLFALP 176
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Db
        301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESP-----VGKEDRVVSPEKTMDIFNEMQ 355
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            Db
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Qy Db		MSVVAPVREEYADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGK 415
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Db	477	
Qу	536	CESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEA 595
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Qy	716	SPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHK-EERLSASPQELGKPY 774
Db	717	SPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVTQHKHKERLSASPQEVGKPY 776
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Db	777	LESFQPNLHITKDAASNEIPTLTKKETISLQMEEFNTAIYSNDDLLSSKEDKMKESETFS 836
Qу	835	DSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSF 894
Db	837	DSSPIEIIDEFPTFVSAKDDSPKEYTDLEVSNKSEIANVQSGANSLPCSELPCDLSF 893
Qy	895	KNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPS 954
Db	894	KNTYPKDEAHVSDEFSKSRSSVSKVPLLLPNVSALESQIEMGNIVKPKVLTKEAEEKLPS 953
Qу	955	DTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVV-FGASLFLLLSLTVFSIVSVTAY 1013
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Db	1074	IKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQAQI 1133
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Db	1134	DHYLGLANKSVKDAMAKIQAKIPGLKRKAE 1163

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    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DE
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GN
    RTN4.
OS
    Mus musculus (Mouse).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
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    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae: Mus.
OX
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RΡ
    STRAIN=129/SvcJ7, and 129SvcJ7;
RC
    Oertle T., van der Putten H., Schwab M.E.;
RA
RT
    "Genomic Structure and Functional Characterization of the Promoter
    Structures of Human and Mouse Nogo/Rtn-4.";
RT
RL
    Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
RN
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RΡ
RC
    STRAIN=129/SvcJ7, and 129SvcJ7;
RA
    Oertle T., Schwab M.E.;
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
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RΡ
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    Van der Putten H.;
RA
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
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RN
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RA
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                       :
                                                 Db
          3 PPLAGGGQKGGAASEAWVPSLFVGVSGSTCTAAKSLVPIPARSSRLSAARNETLFALPAA 62
Qу
         183 SEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGN 242
             Db
         63 SEPVIPSSAEKIMDLKEQPGNTVSSGQEDFPSVLFETAASLPSLSPLSTVSFKEHGYLGN 122
         243 LSAVSSSEGTIEETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAI 302
Qу
            123 LSAVASTEGTIEETLNEASRELPERATNPFVNRESAEFSVLEYSEMGSSFNGSPKGESAM 182
Db
         303 LVENTKEEVIVRSKDKEDLVCSAALHSPQESP-----VGKEDRVVSPEKTMDIFNEMQMS 357
QУ
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Db	183	LVENTKEEVIVRSKDKEDLVCSAALHNPQESPATLTKVVKEDGVMSPEKTMDIFNEMKMS	242
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Qу	418	EGRNEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEE	477
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Qу	478	RKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACE	537
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Qy	538	SELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPL	597
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Qy	598	NSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKALGTKEGIKEP	657
Db	483	NSLLPSTGASVAQPSASPLEVPSPVSYDGIKLEPENPPPYEEAMSVALKTSDSKEEIKEP	542
Qy	658	ESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSP	717
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Qу	718	ESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHK-EERLSASPQELGKPYLE	776
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Qy	777	SFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDS	836
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Qу	837	SPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKN	896
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Qy	957	EKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIAL	1016
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Qy	1017	ALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKE	1076
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GN
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OX
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          Oertle T., van der Putten H., Schwab M.E.;
RA
RT
          "Genomic Structure and Functional Characterization of the Promoter
RT
          Structures of Human and Mouse Nogo/Rtn-4.";
RL
          Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
RN
          [2]
RΡ
          SEQUENCE FROM N.A.
          Oertle T., Schwab M.E.;
          Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
RN
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RP
          SEQUENCE FROM N.A.
RA
          Van der Putten H.;
          Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
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RΡ
          SEQUENCE FROM N.A.
RC
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          Oertle T., van der Putten H., Schwab M.E.;
RT
          "Genomic Structure and Functional Characterization of the Promoter
RT
          Structures of Human and Mouse Nogo/Rtn-4.";
RT.
          J. Mol. Biol. 325:299-323(2003).
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          EMBL; AY102285; AAM64244.1; -.
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QУ
                    255 ETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSFKGSPKGESAILVENTKEEVIVR 314
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Db
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QY		315	SKDRED-LVCSAALHSPQESPVGREDRVVSPERIMDIFNEMQMSVVAPVREEYAD	368
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Qу		369	FKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGRNEDA	424
Db		181	FKPFERVWEVKDSKEDS-DMLAAGGKIESNLESKVDKKCFADSLEQTNHEKDSESSNDDT	239
Qу		425	SFPSTPEPVKDSSRAYITCASFT-SATESTTANTFPLLEDHTSENKTDEKKIEERKAQII	483
Db		240	SFPSTPEGIKDRSGAYITCAPFNPAATESIATNIFPLLGDPTSENKTDEKKIEEKKAQIV	299
Qу		484	TEK-TSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTPDLVQEACESELNE	542
Db		300	TEKNTSTKTSNPFLVAAQDSETDYVTTDNLTKVTEEVVANMPEGLTPDLVQEACESELNE	359
Qу		543	ATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLP	602
Db		360	VTGTKIAYETKMDLVQTSEVMQESLYPAAQLCPSFEESEATPSPVLPDIVMEAPLNSAVP	419
Qу		603	SAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVALKAL-GTKEGIKEPESFN	661
Db		420	SAGASVIQPSSSPLEA-SSVNYESIKHEPENPPPYEEAMSVSLKKVSGIKEEIKEPENIN	478
Qу		662	AAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEP	721
Db		479	AALQETEAPYISIACDLIKETKLSAEPAPDFSDYSEMAKVEQPVPDHSELVEDSSPDSEP	538
Qу		722	VDLFSDDSIPEVPQTQEEAVMLMKESLTEVS-ETVAQHK-EERLSASPQELGKPYLESFQ	779
Db		539	VDLFSDDSIPDVPQKQDETVMLVKESLTETSFESMIEYENKEKLSALPPEGGKPYLESFK	598
Qу		780	PNLHSTKDA-ASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSP: : ::::::::::::::::::::::::	838
Db		599	LSLDNTKDTLLPDEVSTLSKKEKIPLQMEELSTAVYSNDDLFISKEAQIRETETFSDSSP	658
Qy		839	IEIIDEFPTFVSAKDDS-PKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNI	897
Db		659	IEIIDEFPTLISSKTDSFSKLAREYTDLEVSHKSEIANAPDGAGSLPCTELPHDLSLKNI	718
Qy		898	YPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSD	955
Db	-())	719	QPKVEEKISFSDDFSKNGSATSKVLLLPPDVSALATQAEIESIVKPKVLVKEAEKKLPSD	778
Qу		956	TEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIA	1015
Db		779	TEKEDRSPSAIFSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIA	838
Qy	1	.016	LALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIK	1075
Db		839	LALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNCTIK	898
Qу	1	.076	ELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDH	1135
Db		899	ELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSVPVIYERHQAQIDH	958

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    01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DT
DE
    Hypothetical protein.
GN
    RTN4.
    Mus musculus (Mouse).
OS
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC.
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
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RN
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RP
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RΑ
    Strausberg R.;
    Submitted (JUN-2002) to the EMBL/GenBank/DDBJ databases.
RL
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DR
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DR
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DR
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          1 MPEGLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESIYPTAQLCPSFEEAEA 60
        583 TPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMN 642
Qу
            Db
         61 TPSPVLPDIVMEAPLNSLLPSTGASVAOPSASPLEVPSPVSYDGIKLEPENPPPYEEAMS 120
        643 VALKALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFE 702
Qу
                   ┫╏╶┩┦╏╏╃╟┟╬╫╴╊╟┈╏<del>╒</del>╬╟╏╏┩╏╏╏┩┇╏╏┇╬┩╏┦╏┩╒╏╟╟┇┦┩┋╏┦╏╂
        121 VALKTSDAKEEIKEPESFNAAAQEAEAPYISIACDLIKETKLSTEPSPEFSNYSEIAKFE 180
Db
        703 KSVPEHAELVEDSSPESEPVDLFSDDSIPEVPOTOEEAVMLMKESLTEVSETVAOHK-EE 761
Qу
            181 KSVPDHCELVDDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVTQHKHKE 240
Db
        762 RLSASPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLS 821
Qу
            Db
        241 RLSASPQEVGKPYLESFQPNLHITKDAASNEIPTLTKKETISLQMEEFNTAIYSNDDLLS 300
        822 SKEDKIKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGAD 881
Qу
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301 SKEDKMKESETFSDSSPIEIIDEFPTFVSAKDDSP---KEYTDLEVSNKSEIANVOSGAN 357

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         942 KSLTKEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLS 1001
Qу
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        1002 LTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVOK 1061
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        1062 YSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFS 1121
Ov
            Db
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AC
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    01-MAR-2003 (TrEMBLrel. 23, Created)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
DT
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DE
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GN
    RTN4.
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
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RΡ
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RC
    STRAIN=129/SvcJ7, and 129SvcJ7;
    Oertle T., van der Putten H., Schwab M.E.;
    "Genomic Structure and Functional Characterization of the Promoter
    Structures of Human and Mouse Nogo/Rtn-4.";
RT
RL
    Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
RN
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RP
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RC
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    Oertle T., Schwab M.E.;
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RN
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RΡ
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RL
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RN
RΡ
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    STRAIN=129SvcJ7;
RC:
RA
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RL
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
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 Best Local Similarity
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 Matches 367; Conservative
                          2; Mismatches 6; Indels 788; Gaps
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           60 PAAGLSAAPVPP-AAAPLLDFSSDSVPPAPRGPLPAAPPTAPERQPSWERSPAASAPSLP 118
Db
QУ
        121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
           Db
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       181 AASEPVIPSSAEKIMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYL 240
Qу
           Db
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Qу
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Db
       301 AILVENTKEEVIVRSKDKEDLVCSAALHSPQESPVGKEDRVVSPEKTMDIFNEMOMSVVA 360
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Db
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Qу
Db
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Qу
       421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480
Db
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Qy
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Db
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Db
Qу
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Db
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Oy
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Db
Qу
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                         Db
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        1021 VTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRL 1080
Qу
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Db
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            Db
        293 FLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQAQIDHYLGLA 352
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            Db
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RESULT 7
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ID
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AC
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DT
    01-MAR-2003 (TrEMBLrel. 23, Created)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DΤ
DE
    RTN4.
GN
    RTN4.
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
    NCBI TaxID=10090;
OX
RN
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    SEQUENCE FROM N.A.
RP
RC
    STRAIN=129/SvcJ7, and 129SvcJ7;
RA
    Oertle T., van der Putten H., Schwab M.E.;
RT
    "Genomic Structure and Functional Characterization of the Promoter
RТ
    Structures of Human and Mouse Nogo/Rtn-4.";
RL
    Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
RN
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=129/SvcJ7, and 129SvcJ7;
RA
    Oertle T., Schwab M.E.;
RL
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
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RN
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RP
    SEQUENCE FROM N.A.
RC
    STRAIN=129/SvcJ7;
RA
    Van der Putten H.;
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
RN
RΡ
    SEQUENCE FROM N.A.
RC
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    Van der Putten H., Mir A.;
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
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    EMBL; AY102286; AAM73508.1; -.
DR
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 Best Local Similarity 29.9%; Pred. No. 2.5e-59;
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       121 PAAAVLPSKLPEDDEPPARPPPPPPAGASPLAEPAAPPSTPAAPKRRGSGSVDETLFALP 180
Qу
           Db
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Db
       169 ----- 168
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Qу
Db
       169 ----- 168
Qу
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Dh
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Qу
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Db
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Db
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Qу
       169 ----- 168
Db
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Db
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Qу
Db
Qу
        721 PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAOHKEERLSASPOELGKPYLESFOP 780
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Db
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        169 ----- 168
Db
Qу
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Db
        169 ----- 168
Qу
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        169 ----- 168
Db
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Qу
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AC
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DT
    01-OCT-2002 (TrEMBLrel. 22, Created)
DT
    01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DT
DE
   Nogo-B.
GN
   RTN4.
OS
   Mus musculus (Mouse).
   Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
OC
   Mammalia; Eutheria; Rodentia; Sciuroqnathi; Muridae; Murinae; Mus.
OX
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RN
    [1]
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RP

SEQUENCE FROM N.A.

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RC
    STRAIN=BALB/C;
RA
    Jin W., Li R., Long M., Shen J., Ju G.;
    "Cloning and expression of the mouse Nogo-B protein.";
RT
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RL
    EMBL; AY114153; AAM77069.1; -.
DR
    MGD; MGI:1915835; Rtn4.
DR
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    Pfam; PF02453; Reticulon; 1.
DR
DR
    PROSITE; PS50845; RETICULON; 1.
SO
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 Best Local Similarity 29.9%; Pred. No. 8.4e-59;
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Db
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Dh
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Qу
Db
Qу
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Qу
Db
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QУ
       169 ----- 168
Db
Qу
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Qу	721	PVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSASPQELGKPYLESFQP	780
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Qу	781	NLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIE	840
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Qy	841	IIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPCLELPCDLSFKNIYPK	900
Db	169		168
Qу	901	DEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKLPSDTEKED	960
Db	169		168
Qу	961	RSLSAVLSAELSKTSVVDLLYWRDIKKTGVV-FGASLFLLLSLTVFSIVSVTAYIALALL	1019
Db	169	VVDLLYWRDIKKTGVVYFGASLFLLLSLTVFSIVSVTAYIALALL	213
QУ	1020	SVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRR	1079
Db	214	SVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRR	273
Qу	1080	LFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGL	1139
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Qу	1140	ANKSVKDAMAKIQAKIPGLKRKAD 1163	
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Q96E ID AC DT DT DT DE GN OS	Q96B16; Q96B16; 01-DEC-20 01-DEC-20 01-MAR-20 Hypothet: RTN4. Homo sap:	PRELIMINARY; PRT; 392 AA. 001 (TrEMBLrel. 19, Created) 001 (TrEMBLrel. 19, Last sequence update) 003 (TrEMBLrel. 23, Last annotation update) ical protein (RTN4).	
OC OX RN RP RC RA RL	Mammalia NCBI_Taxi [1] SEQUENCE TISSUE=Ki Strausber	FROM N.A.	

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RN
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RP
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RA
    Oertle T., van der Putten H., Schwab M.E.;
RT
    "Genomic Structure and Functional Characterization of the Promoter
RT
    Structures of Human and Mouse Nogo/Rtn-4.";
RL
    Submitted (OCT-2002) to the EMBL/GenBank/DDBJ databases.
RN
    [3]
RP
    SEQUENCE FROM N.A.
RA
    Oertle T., Schwab M.E.;
RL
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RN
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RP
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RA
    Van der Putten H.;
RT.
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
RN
    [5]
RP
    SEQUENCE FROM N.A.
RX
    MEDLINE=22376540; PubMed=12488097;
RA
    Oertle T., van der Putten H., Schwab M.E.;
RТ
    "Genomic Structure and Functional Characterization of the Promoter
RT
    Structures of Human and Mouse Nogo/Rtn-4.";
RL
    J. Mol. Biol. 325:299-323(2003).
DR
    EMBL; BC016165; AAH16165.1; -.
DR
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DR
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DR
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DR
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DR
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KW
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 Best Local Similarity
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Db
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Db
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Qу
                 Db
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Qу
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Qу	466	${\tt SENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPE}$	525
Db	205		204
Qу	526	GLTPDLVQEACESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAQLCPSFEEAEATPS	585
Db	205		204
Qу	586	PVLPDIVMEAPLNSLLPSAGASVVQPSVSPLEAPPPVSYDSIKLEPENPPPYEEAMNVAL	645
Db	205		204
Qу	646	KALGTKEGIKEPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSV	705
Db	205		204
Qу	706	PEHAELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHKEERLSA	765
Db	205		204
Qу	766	${\tt SPQELGKPYLESFQPNLHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKED}$	825
Db	205		204
Qy	826	KIKESETFSDSSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIQSGADSLPC	885
Db	205		204
Qy	886	LELPCDLSFKNIYPKDEVHVSDEFSENRSSVSKASISPSNVSALEPQTEMGSIVKSKSLT	945
Db	205		204
Qy	946	KEAEKKLPSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVF	1005
Db	205		234
Qy	1006	SIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNS	1065
Db	235		294
Qy	1066	ALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVI	1125
Db	295	ALGHVNCTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSVPVI	354
Qy .	1126	YERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163	
Db	355	: : YERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 392	

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RESULT 10
Q9GM33
ID
               PRELIMINARY;
    Q9GM33
                                PRT; 179 AA.
AC
    09GM33;
    01-MAR-2001 (TrEMBLrel. 16, Created)
DТ
DT
    01-MAR-2001 (TrEMBLrel. 16, Last sequence update)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
    Hypothetical 19.9 kDa protein.
DE
    Macaca fascicularis (Crab eating macaque) (Cynomolgus monkey).
OS
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Mammalia; Eutheria; Primates; Catarrhini; Cercopithecidae;
OC
OC
    Cercopithecinae; Macaca.
    NCBI TaxID=9541;
OX
RN
    [1]
    SEQUENCE FROM N.A.
RΡ
RC
    TISSUE=Brain:
RA
    Osada N., Hida M., Kusuda J., Tanuma R., Iseki K., Hirai M., Terao K.,
RA
    Suzuki Y., Sugano S., Hashimoto K.;
RT
    "Isolation of full-length cDNA clones from macaque brain cDNA
    libraries.";
RT
RL
    Submitted (OCT-2000) to the EMBL/GenBank/DDBJ databases.
DR
    EMBL; AB049853; BAB16739.1; -.
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
DR
    PROSITE; PS50845; RETICULON; 1.
KW
    Hypothetical protein.
SQ
    SEQUENCE
             179 AA; 19949 MW; 5F8CD4383FEE9E02 CRC64;
  Query Match
                        14.4%; Score 844; DB 6; Length 179;
 Best Local Similarity
                      95.5%; Pred. No. 8.5e-36;
 Matches 171; Conservative 6; Mismatches
                                              2; Indels
                                                            0; Gaps
                                                                       0;
Qу
         985 IKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVIQAIQKSDEGHPF 1044
             Db
           1 MKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYRGVIQAIQKSDEGHPF 60
        1045 RAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKFAVLMWVFTYVGA 1104
Qу
             Db
          61 RAYLESEVAISEELVQKYSNSALGHVNCTIKELRRLFLVDDLVDSLRFAVLMWVFTYVGA 120
        1105 LFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
Qу
             Dh
         121 LFNGLTLLILALISLFSVPVIYERHQAQIDHYLGLANKNVKDAMAKIQAKIPGLKRKAE 179
RESULT 11
Q8K4S4
ID
    Q8K4S4
               PRELIMINARY;
                                PRT;
                                       780 AA.
AC
    Q8K4S4;
DΤ
    01-OCT-2002 (TrEMBLrel. 22, Created)
DT
    01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
    Reticulon 1A.
GN
    RTN1 OR RTN-1A.
OS
    Mus musculus (Mouse).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
```

```
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
    SEQUENCE FROM N.A.
RΡ
RC
    STRAIN=ICR; TISSUE=Brain;
RA
    Hirata T., Nomura T., Takagi Y., Sato Y., Tomioka N., Fujisawa H.,
RA
RT
    "Mosaic development of the olfactory cortex with Pax6-dependent and
RТ
    -independent components.";
    Submitted (NOV-2001) to the EMBL/GenBank/DDBJ databases.
RL
    EMBL; AB074899; BAB96551.1; -.
DR
DR
    MGD; MGI:1933947; Rtn1.
DR
    InterPro; IPR001951; Histone H4.
    InterPro; IPR003388; Reticulon.
DR
DR
    Pfam; PF02453; Reticulon; 1.
    PROSITE; PS00047; HISTONE H4; 1.
DR
DR
    PROSITE; PS50845; RETICULON; 1.
    SEQUENCE 780 AA; 83504 MW; 545F5638C576A069 CRC64;
SQ
                     13.5%; Score 792; DB 11; Length 780;
 Query Match
 Best Local Similarity 32.5%; Pred. No. 3e-32;
 Matches 242; Conservative 105; Mismatches 260; Indels 138; Gaps
        517 EAAVSNMPEGLTPDLVQEACESELNEATG-----TKIAYETKVDLVQTSEAIQ-ESLY 568
Qу
           76 ETASTGM--AAVPDALDHSPSSTLKDGEGACYTSLISDVCYPPREDSAYFTGILQKENGH 133
Db
        569 PTAQLCPSFEEAEATPSPVLPDIVMEAP---LNS-----LLPSAGASVVQPSVSPLEAPP 620
Qу
                134 ITTSESP--EEPE-TPGPSLPEVPGMEPQGLLSSDSGIEMTPAESTEVNKILADPLDQMK 190
Db
        621 PVSYDSIKLE-----PENPPPYEEAMNVALKALGTK----EGIKEPESFNAAV---- 664
Qу
             191 AEAYKYIDITRPQEAKGQEEQHPGLEDKDLDFKDKGTEVSTKAEGVRAPNQ-PAPVEGKL 249
Db
        665 -----QETEAPYISIACDLIKETKLSTEPSPDFSNYSEI----AKFEKSVPEHAELV 712
Qу
                 Db
        250 IKDHLFEESTFAPYIDELSD--EOHRVSLVTAPVKITLTEIEPPLMTATOETIPEKODLC 307
        713 EDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAQHK-----EERLSASP 767
Qу
              308 LKPSPDTVPTVTVSE-----PEDDSPGSVTPPSSGTEPSAAESQGKGSVSEDELIAAIK 361
Db
        768 QELGKPYLESFQPN-----LHSTKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSS 822
Qу
           : | : | : :|:
        362 EAKGLSYETTESPRPVGQVADKPKTKTRSGLPTIPS-----PLDQEASS 405
Db
        823 KEDKIKESETFSDSSPIEIIDEFPT-----FVSAKDDSPKLAK-EYTDLEVSDKSEIAN 875
Qу
               :|: | ::|:
        406 AESGDSEIELVSE-DPMASEDALPSGYVSFGHVSGPPPSPASPSIQYSILREEREAEL-- 462
Db
        876 IOSGADSLPCLELPCDL-SFKNIYPKDEVH------VSDEFSENRSSVSKASISPSNVS 927
Qу
                                     463 ----DSELIIE-SCDASSASEESPKREQDSPPMKPGALDAIREETGSRATEERAPSHQG 516
Db
        928 ALEPOTEMGSIVKSKSLTKEAEKKL-----PSDTEKEDRSLSAVLSAE----- 970
Ov
            :|| || :::: || |:::
```

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Db
        517 PVEPD-PMLSFAPAAALQSRPEPSSGDGASVPEPPRSQQQKPEEEAVSSSOSPTATEIPG 575
ОУ
        971 -----LSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALAL 1018
                      576 PLGSGLMPPLPFFNKQKAIDLLYWRDIKQTGIVFGSFLLLLFSLTQFSVVSVVAYLALAA 635
Db
       1019 LSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELR 1078
Qу
            636 LSATISFRIYKSVLQAVQKTDEGHPFKAYLELEITLSQEQIQKYTDCLQLYVNSTLKELR 695
Db
       1079 RLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLG 1138
Qу
            Db
        696 RLFLVQDLVDSLKFAVLMWLLTYVGALFNGLTLLLMAVVSMFTLPVVYVKHQAQVDQYLG 755
       1139 LANKSVKDAMAKIQAKIPGLKRKAD 1163
QУ
            Db
        756 LVRTHINTVVAKIQAKIPGAKRHAE 780
RESULT 12
Q8K0T0
ID
    O8KOTO
              PRELIMINARY;
                            PRT;
                                    780 AA.
AC
    Q8K0T0;
DT
    01-OCT-2002 (TrEMBLrel. .22, Created)
    01-OCT-2002 (TrEMBLrel. 22, Last sequence update)
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DT
DE
    Similar to reticulon 1.
GN
    RTN1.
OS
    Mus musculus (Mouse).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciuroqnathi; Muridae; Murinae; Mus.
OX
    NCBI TaxID=10090;
RN
    [1]
RP
    SEQUENCE FROM N.A.
RC
    TISSUE=Retina;
RA
    Strausberg R.;
RT_1
    Submitted (MAY-2002) to the EMBL/GenBank/DDBJ databases.
DR
    EMBL; BC030455; AAH30455.1; -.
DR
    MGD; MGI:1933947; Rtnl.
DR
    InterPro; IPR001951; Histone H4.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
DR
    PROSITE; PS00047; HISTONE H4; 1.
DR
    PROSITE; PS50845; RETICULON; 1.
SO
    SEQUENCE 780 AA; 83572 MW; 29B47A58FC2F2027 CRC64;
 Query Match
                      13.5%; Score 788; DB 11; Length 780;
 Best Local Similarity 32.4%; Pred. No. 4.9e-32;
 Matches 243; Conservative 104; Mismatches 256; Indels 146; Gaps
                                                                 28:
QУ
        517 EAAVSNMPEGLTPDLVQEACESELNEATG-----TKIAYETKVDLVQTSEAIQ-ESLY 568
            Db
         76 ETASTGM--AAVPDALDHSPSSTLKDGEGACYTSLISDVCYPPREDSAYFTGILOKENGH 133
        569 PTAQLCPSFEEAEATPSPVLPDIVMEAP---LNS-----LLPSAGASVVQPSVSPLEAPP 620
Qу
            134 ITTSESP--EEPE-TPGPSLPEVPGMEPQGLLSSDSGIEMTPAESTEVNKILADPLDQMK 190
Db
```

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Qу
        621 PVSYDSI-----KLEPENPPPYEEA------MNVALKALGTKEGIKEPESFNAAV 664
                      |: ||
                                                Db
        191 AEAYKYIDITRPQEAKGQEEQHPGLEDKDLDFKDKDTEVSTKA----EGVRAPNQ-PAPV 245
        665 ------QETEAPYISIACDLIKETKLSTEPSPDFSNYSEI-----AKFEKSVPEH 708
Qу
                    : | ||||
                            | :: ::| :| :||
        246 EGKLIKDHLFEESTFAPYIDELSD--EQHRVSLVTAPVKITLTEIEPPLMTATQETIPEK 303
Db
        709 AELVEDSSPESEPVDLFSDDSIPEVPQTQEEAVMLMKESLTEVSETVAOHK----EERL 763
Qу
               304 ODLCLKPSPDTVPTVTVSE-----PEDDSPGSVTPPSSGTEPSAAESQGKGSVSEDELI 357
Db
        764 SASPOELGKPYLESFOPN-----LHSTKDAASNDIPTLTKKEKISLOMEEFNTAIYSNDD 818
Qу
           358 AAIKEAKGLSYETTESPRPVGQVADKPKTKTRSGLPTIPS-----PLDQ 401
Db
        819 LLSSKEDKIKESETFSDSSPIEIIDEFPT----FVSAKDDSPKLAK-EYTDLEVSDKS 871
Qу
             Db
        402 EASSAESGDSEIELVSE-DPMASEDALPSGYVSFGHVSGPPPSPASPSIOYSILREEREA 460
Qу
        872 EIANIQSGADSLPCLELPCDL-SFKNIYPKDEVH------VSDEFSENRSSVSKASISP 923
           | | : :|
        461 EL-----DSELIIE-SCDASSASEESPKREODSPPMKPGALDAIREETGSRATEERAP 512
Db
        924 SNVSALEPQTEMGSIVKSKSLTKEAEKKL-----PSDTEKEDRSLSAVLSAE---- 970
Qу
           513 SHQGPVEPD-PMLSFAPAAALQSRPEPSSGDGASVPEPPRSQQQKPEEEAVSSSQSPTAT 571
Db
        971 -----LSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYI 1014
Qу
                         Db
        572 EIPGPLGSGLMPPLPFFNKQKAIDLLYWRDIKOTGIVFGSFLLLLFSLTOFSVVSVVAYL 631
Qу
       1015 ALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVOKYSNSALGHVNSTI 1074
           632 ALAALSATISFRIYKSVLQAVQKTDEGHPFKAYLELEITLSQEQIQKYTDCLQLYVNSTL 691
Db
       1075 KELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHOVOID 1134
Qу
           692 KELRRLFLVQDLVDSLKFAVLMWLLTYVGALFNGLTLLLMAVVSMFTLPVVYVKHQAQVD 751
Dh
Qу
       1135 HYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
            Db
        752 QYLGLVRTHINTVVAKIQAKIPGAKRHAE 780
RESULT 13
090638
ID
    Q90638
             PRELIMINARY;
                          PRT; 760 AA.
AC
    Q90638;
DT
    01-NOV-1996 (TrEMBLrel. 01, Created)
DT
    01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DE
    ChS-Rex-b.
OS
    Gallus gallus (Chicken).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
    Archosauria; Aves; Neognathae; Galliformes; Phasianidae; Phasianinae;
```

OC.

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OC
    Gallus.
    NCBI TaxID=9031;
OX
RN
RP
    SEQUENCE FROM N.A.
RC
    TISSUE=Brain;
RX
    MEDLINE=96386034; PubMed=8793864;
    Baka I.D., Ninkina N.N., Pinon L.G., Adu J., Davies A.M.,
RA
RA
    Georgiev G.P., Buchman V.L.;
RT
    "Intracellular compartmentalization of two differentially spliced s-
    rex/NSP mRNAs in neurons.";
RT
    Mol. Cell. Neurosci. 7:289-303(1996).
RL
RN
RΡ
    SEQUENCE FROM N.A.
RC
    TISSUE=Brain;
RX
    MEDLINE=97183663;
RA
    Ninkina N.N., Baka I.D., Buchman V.L.;
RT
    "Rat and chicken s-rex/NSP mRNA: nucleotide sequence of main
    transcripts and expression of splice variants in rat tissues.";
RT
    Gene 184:205-210(1997).
RL
DR
    EMBL; U17606; AAC60075.1; -.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
    PROSITE; PS50845; RETICULON; 1.
SO
             760 AA; 82502 MW; 465C1B429F799D5C CRC64;
    SEQUENCE
 Query Match 13.0%; Score 762.5; DB 13; Length 760; Best Local Similarity 27.1%; Pred. No. 9.3e-31;
 Matches 285; Conservative 124; Mismatches 319; Indels 323; Gaps
                                                                  34;
         142 PPPP----AG-ASPLAEPAAPPSTPAAPKRRGSGSVDETLFALPAASE--PVIPSSAEK 193
Qу
                   4 PPDPQDLLLAGTAERWAAAGADEYAAGAALRDGDGAQQREQLAFGSAREHPPVAMATA-- 61
Db
        194 IMDLMEQPGNTVSSGQEDFPSVLLETAASLPSLSPLSTVSFKEHGYLGNLSAVSSSEGTI 253
Qу
                  62 -----SPGVTASSRLFDYGSSSANGADSSFYTSLISDVHY----- 96
Db
         254 EETLNEASKELPERATNPFVNRDLAEFSELEYSEMGSSF--KGSPKGESAILVENTKEEV 311
Qу
                         ----TTP---RDNTYFTGV-YQQENSPIPCSGSTEGFN----- 126
Db
        312 IVRSKDKEDLVCSAALHSPQESPVGKEDRVV-----SPEKTMDIFNEMQMSVVAPVREEY 366
Qу
                        Db
        127 -----ALGHPVQDVTGFESRGLFSLDSGIEMTPAESAEVDKSLTDPMKVEG 172
        367 ADFKPFEQAWEVKDTYEGSRDVLAARANVESKVDRKCLEDSLEQKSLGKDSEGR----- 420
QУ
              Db
        173 YKYMDISRPEEIK--YOEKHD-----PDSEDESPDLIDEYRGTPIGSGHAAEPORTTA 223
        421 NEDASFPSTPEPVKDSSRAYITCASFTSATESTTANTFPLLEDHTSENKTDEKKIEERKA 480
Qу
                Db
        224 SEAIKAPKEODPLEDKSFRDOHNASVVTAPVKITLTETPGAREATS-----KEA 272
        481 QIITEKTSPKTSNPFLVAVQDSEADYVTTDTLSKVTEAAVSNMPEGLTP----DLVQEAC 536
QУ
             Db
        273 SVTQPKSGLKPSHEVVPTVMVSEPE---DDSPGSVTPPSSGTEPSGSESQGKGSLSEDEL 329
```

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537 ESELNEATGTKIAYETKVDLVQTSEAIQESLYPTAOLCPSFEEAEATPSPVLPDIVMEAP 596
Qу
            330 ISAIKEAKG--FSFETSE--VOOSPAV------SAEKOEOKMKPGRPAV---- 368
Db
        597 LNSLLPSAGASVVQPSVSPLEAPPPVSYDS-IKLEPENPPPYEEAMNVALKALGTKEGIK 655
Qу
                      369 -----PSPLDNEASSAESGDSETELVSEDPLAAEEVLHSNYMTFSHIGG-- 412
Db
        656 EPESFNAAVQETEAPYISIACDLIKETKLSTEPSPDFSNYSEIAKFEKSVPEHAELVEDS 715
QУ
                                   413 PPPS------PASPSIQYS----ILREEREAELDSELIIES 443
Dh
QУ
        716 SPESEPVDLFSDDSIPEVPOTOEEAVMLMKESLTEVSETVAOHKEERLSASPOELGKPYL 775
                     444 CDAS-----SASEESPKREODSPLMKPMVMDI---IKEENSSRASASDYEASK--- 488
Db
QУ
        776 ESFQPNLHSTKDAASNDIPTLTKKEKISLOMEEFNTAIYSNDDLLSSKEDKIKESETFSD 835
                   | :: |
                                                  : : : ::
                           -----RENLADSASYLK 507
Db
               ----TTESRMN---
Qу
        836 SSPIEIIDEFPTFVSAKDDSPKLAKEYTDLEVSDKSEIANIOSGADSLPCLELPCDLSFK 895
           | | |
                | |
                         : | |,: : |
        508 SS------ FV---- APKVSSE---- 518
Db
        896 NIYPKDEVHVSDEFSEN---RSSVSKASISPSNVSALEPQTEMGSIVKSKSLTKEAEKKL 952
Qу
             Db
        519 --- PPTSAVSTEELKERIILKKPIEETVVNOSKVS---
        953 PSDTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTA 1012
Qу
           Db
        559 P-----LALPLLPFLNKQKAINLLYWRDIKQTGIVFGSLLLLLFSLTQFSVVSVVA 609
       1013 YIALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNS 1072
Qу
            610 YLALAGLSATISFRIYKSVLQAVQKTDEGHPFKAYLDMEMNLSQDQIQKYTDCLQLYVNS 669
Db
Qу
       1073 TIKELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHOVO 1132
           Db
        670 TVKELRRLFLVQDLVDSLKFAVLMWLLTYVGALFNGLTLLIMAVVSMFTLPVVYDKYOAO 729
       1133 IDHYLGLANKSVKDAMAKIQAKIPGLKRKAD 1163
Qу
                  : :|||||||||
Db
        730 IDQYLGLVRTHINTVVAKIQAKIPGAKRKAE 760
RESULT 14
Q8CCU2
                           PRT; 643 AA.
ID
    Q8CCU2
              PRELIMINARY;
AC
    Q8CCU2;
DT
    01-MAR-2003 (TrEMBLrel. 23, Created)
    01-MAR-2003 (TrEMBLrel. 23, Last sequence update)
DT
DT
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DE
    Reticulon 3.
OS
    Mus musculus (Mouse).
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
OC.
OX
    NCBI TaxID=10090;
```

```
RN
    [1]
RP
    SEQUENCE FROM N.A.
RC
    STRAIN=C57BL/6J; TISSUE=Medulla oblongata;
    MEDLINE=22354683; PubMed=12466851;
RX
RA
    The FANTOM Consortium,
    the RIKEN Genome Exploration Research Group Phase I & II Team;
RT
    "Analysis of the mouse transcriptome based on functional annotation of
RT
    60,770 full-length cDNAs.";
RL
    Nature 420:563-573(2002).
DR
    EMBL; AK032109; BAC27708.1; -.
    SEQUENCE 643 AA; 68769 MW; 15E3335B280533F8 CRC64;
SO
 Query Match
                    12.0%; Score 700; DB 11; Length 643;
 Best Local Similarity 30.3%; Pred. No. 1.1e-27;
 Matches 227; Conservative 109; Mismatches 234; Indels 180; Gaps
QУ
        442 TCA-SFTSATESTTANTFPLLEDHTSENKTDEKKIEERKAQIITEKTSPKTSNPFLVAVO 500
           ;|| || |:: | : : | : |: |:
Db
         46 SCADSFVSSSSSQPVSIFSTSQGDWTEAFTEGKPVR-----
Qу
        501 DSEADYVTTDTLSKVTEAAVSNMP-----EGLTPDLVQEACESELNEATGTKIAYETKVD 555
                   Db
              -----DYLSSTKEAGGNGVPGSSQLHSELPGSMPEKWVSGSGAAT-----VEVT 125
        556 LVQTSEAIQESLYPTAQLCPSFEEAEATPSPVLPDIVMEAPLNSLLPSAGASVVQPSVSP 615
Qу
           126 LPNLRGAWPNSVMGEVTEVDSSGESDDT---VIEDIT-EKP--DSLPSAAAKTSEREIK- 178
Db
        616 LEAPPPVSYDSIKLE--PENPPPYEEAMNVALKALGTKEGIKEPESFNAAVQE----TEA 669
QУ
            Db
        179 -ETP---SRETVRSEMCENSEQPQAQPETPTQKSLEGEVASQVPNTLNEVTPEKLDMTNN 234
        670 PYISIAC--DLIKETKLS-TEPSPDFSNYSEIAKFEKSVPEHAELVEDSSPESEPVDLFS 726
Qу
           235 PKVCSAAPPSVLNETGFSLTVPAS------AKLESLLGKYVEDTDGSSPE----- 278
Db
QУ
        727 DDSIPEVPQTQEEAVMLMKES--LTEVSETVAQHKEERLSASPQELGKPYLESFQPNLHS 784
            Db
        279 -DLMAVLTGAEEKGIVDKEEGDVLEAVLEKIADFK----NTLPVEL------LHE 322
        785 TKDAASNDIPTLTKKEKISLQMEEFNTAIYSNDDLLSSKEDKIKESETFSDSSPIEIIDE 844
Qу
                                        Db
                                    ----LSGSETKNIKSKYSEDSR--ETTGG 347
        845 FPTFVSAKDDSPKLAKEYTDLEV-----SDKSEIANIQSGADSLPCLELPCDLSFKN 896
Qу
                   Db
        348 APTM-----SPDLEQEQLTIRAIKELGERQAEKVQDEGISSGG------KLKQ 389
        897 IY-PKDEVHVSDEFSENRSSVSKASIS-PSNVSALEPQTEMGSIVKSKSLTKEAEKKLPS 954
Qу
            Db
        390 TFAPQSGPQSSSDILEHTDVKTGSDLGIPKNPTIIK-NTRIDSI---SSLTK----- 437
        955 DTEKEDRSLSAVLSAELSKTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYI 1014
Qу
            438 -TEMVNKN---VLARLLSDFPVHDLIFWRDVKKTGFVFGTTLIMLLSLAAFSVISVVSYL 493
Db
Qу
       1015 ALALLSVTISFRIYKGVIQAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTI 1074
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Db
        494 ILALLSVTISFRVYKSVIQAVQKSEEGHPFKAYLDVDITLSSEAFHNYMNAAMVHVNKAL 553
       1075 KELRRLFLVDDLVDSLKFAVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQID 1134
Qу
            554 KLIIRLFLVEDLVDSLKLAVFMWLMTYVGAVFNGITLLILAELLVFSVPIVYEKYKTQID 613
Db
       1135 HYLGLANKSVKDAMAKIQAKIPGL-KRKAD 1163
Qу
                   Db
        614 HYVGIARDQTKSIVEKIQAKLPGIAKKKAE 643
RESULT 15
Q9BQ59
ID
    09B059
              PRELIMINARY;
                             PRT:
                                   199 AA.
AC
    Q9BQ59;
DT
    01-JUN-2001 (TrEMBLrel. 17, Created)
    01-JUN-2001 (TrEMBLrel. 17, Last sequence update)
    01-MAR-2003 (TrEMBLrel. 23, Last annotation update)
DT
DE
    Similar to reticulon 1.
OS
    Homo sapiens (Human).
OC
    Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
OC
    Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
    NCBI TaxID=9606;
OX
RN
    [1]
    SEQUENCE FROM N.A.
RP
RC.
    TISSUE=Lung;
RA
    Strausberg R.;
    Submitted (FEB-2001) to the EMBL/GenBank/DDBJ databases.
RL
DR
    EMBL; BC003003; AAH03003.1; -.
DR
    EMBL; BC000314; AAH00314.1; -.
DR
    InterPro; IPR001951; Histone H4.
DR
    InterPro; IPR003388; Reticulon.
DR
    Pfam; PF02453; Reticulon; 1.
    PROSITE; PS00047; HISTONE H4; 1.
DR
DR
    PROSITE; PS50845; RETICULON; 1.
SO
    SEQUENCE 199 AA; 22642 MW; 7CFA44CC568DF6D8 CRC64;
 Query Match
                      11.7%; Score 685; DB 4; Length 199;
 Best Local Similarity 67.0%; Pred. No. 1.2e-27;
 Matches 128; Conservative 32; Mismatches 31; Indels
                                                       0; Gaps
                                                                 0;
        973 KTSVVDLLYWRDIKKTGVVFGASLFLLLSLTVFSIVSVTAYIALALLSVTISFRIYKGVI 1032
Qу
            Db
          9 KSQAIDLLYWRDIKQTGIVFGSFLLLLFSLTQFSVVSVVAYLALAALSATISFRIYKSVL 68
Qу
       1033 QAIQKSDEGHPFRAYLESEVAISEELVQKYSNSALGHVNSTIKELRRLFLVDDLVDSLKF 1092
            Db
         69 QAVQKTDEGHPFKAYLELEITLSQEQIQKYTDCLQFYVNSTLKELRRLFLVQDLVDSLKF 128
QУ
       1093 AVLMWVFTYVGALFNGLTLLILALISLFSIPVIYERHQVQIDHYLGLANKSVKDAMAKIO 1152
            Db
        129 AVLMWLLTYVGALFNGLTLLLMAVVSMFTLPVVYVKHQAQIDQYLGLVRTHINAVVAKIO 188
       1153 AKIPGLKRKAD 1163
Qу
            Db
        189 AKIPGAKRHAE 199
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Search completed: December 19, 2003, 15:36:42

Job time : 51 secs